

HISTOPATHOLOGY OF THE SKIN

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In Memory of My Father

DR ALEXANDER LEVER 1877–1946

1877–1946 My First Teacher in Dermatology



Preface to the Second Edition

Descriptive histopathology is regarded by some as a static science in which nowadays but few concepts change. But this is not so. Many advances have been made in the field of dermatopathology in only the five years that have passed since the appearance of the First Edition of this book. Consequently many changes were necessary for this Second Feltion.

Important advances have been made in recent years in the histo logic diagnosis of the vesicular and the bullous diseases especially of pemphigus Therefore the descriptions of these diseases have been entirely rewritten and brought together into a new separate chapter Because of changes in the concept of the histogenesis of the nevus cell the chapter on nevi and melanomas also has been rewritten The use of the periodic acid Schiff reaction for the demonstration of funct in tissue is an important advance which is taken due notice of in the chapter on fungus diseases. Extensive changes also have been made for instance in the descriptions of kraurosis vulvae the pur puras and verruca Furthermore discussion of the following diseases has been incorporated because recent work has increased the interest of dermatologists in them beryllium granuloma papular myxedema porphyria ochronosis hibernoma and hemangiopericytoma Many changes have been made in the Bibliographies that follow each chapter to order to keep them up to date Fifty six new photomicro graphs have been added and 14 of the old photomicrographs have been replaced by better ones

It is with great regret that I record the death of Dr. Tracy B. Millor) former chief of the Pathology Laboratory at the Massa clinicitis General Hospital. To lim. I owe a great debt of gratitude Dr. Benjamin Castleman, the present chief of the laboratory has dissed and helped me in miny ways and I thank him. I also wish to express my thanks to Mr. Richard W. St. Clair for producing with great skill the new photomicrographs and to Mr. Walter Kahoe Director of the Medical Department of J. B. Lippincoit Company for his many courtesies and his infialing co-operation.

WALTER F LEVER



Preface to the First Edition

This book is based on the courses of dermatopathology which I have been giving in recent years to graduate students of dermatology enrolled at Harvard Medical School and Massychusetts General Hos pital. The book is written primarily for dermatologists. I hope however that it may be useful also to pathologists since dermatopa thology is given little consideration in most textbooks of pathology.

I have attempted to keep this book short Emphriss has been pliced on the essential histologic features. Minor details and rare absertations from the typical histologic picture hive been omitted. I have allotted more space to the cutaneous diseases in which histologic extimination is of diagnostic value thin to those in which the histologic picture is not characteristic. In spite of my striving for brevity. I have discussed the histogenesis of several dermatoses be cause knowledge of the histogenesis often is of great value for the understinding of the pathologic process.

Primarily for the benefit of pathologists who usually are not too familiar with demandologic diseases. I have preceded the histologic discussion of each disease with a short description of the clinical

features

A furly extensive bibliography has been supplied for readers who are interested in obtaining additional information. In the selection of articles for the bibliography preference has been given whenever

possible to those written in English

I wish to express my deep gratitude to Dr Tracy B Mallory and Dr Benjamin Cvistleman of the Pathology Laboratory at the Massa clusetts General Hospital for the truming in pathology, they have given me It has been invaluable to me Their teaching is reflected in this book. Furthermore I wish to than, Mr Richard W St Clair who with great skill and patience produced all the photomicrographs in this book.

WALTER F LEVER



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Introduction

TECHNIC FOR BIOPSY

It is important to select a proper site for biopsy. In most instances histologic examination of a fully developed lesion will give more information than examination of an errly or an involuting lesion. An exception to this rule represents vesicular, bullous and pustular lesions. For their histologic examination a erry early lesion is required otherwise, secondary changes (such as regeneration degeneration or secondary infection) may obscure essential features and make recognition of their mode of formation impossible Generally, it is inadvisable to include normal tissue in the biopsy specimen unless a large specimen is taken or the physician personally super uses the processing of the specimen because improper sectioning by the technician may result in only normal skin being seen in the section. The specimen should include subcutaneous fat, because, in many dermatoses characteristic histologic features are found in the lower dermis or in the subcutaneous fat. If several types of lesions are present and the diagnosis hinges on the histologic findings much time may be saved by taking specimens for biopsy from more than one lesion.

In the author's experience a specimen obtained with a 6 mm biopsy punch nearly always has proved adequate for histologic study. Two sutures are sufficient to close the wound

Before placing the specimen in the routine fixative, which in many hospitals is Zenkers or Helly s solution one should consider which stains are destrable. Some of the special stains can be performed only if the specimen has been fixed in the appropriate fixative. A tabulation of staining methods with the required fixatives is found in Table 1 on page 29.

LIMITATIONS OF HISTOLOGIC DIAGNOSIS

Although histologic study is one of the most valuable means of diagnosis in dermatology, it has its limitations Often no definitive diagnosis can be made. The reason for this is that few dermatoses, aside from the tumors, are associated regularly with a diagnostic his

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Introduction

TECHNIC FOR BIOPSY

It is important to select a proper site for biopsy. In most instances histologic examination of a fully developed lesion will give more information than examination of an early or an involuting lesion An exception to this rule represents vesicular bullous and pustular lesions For their histologic examination a very early lesion is re outed otherwise secondary changes (such as regeneration degen eration or secondary infection) may obscure essential features and make recognition of their mode of formation impossible Generally it is inadvisable to include normal tissue in the biopsy specimen unless a large specimen is taken or the physician personally super vises the processing of the specimen because improper sectioning by the technician may result in only normal skin being seen in the sec tion. The specimen should include subcutaneous fat because in many dermatoses characteristic histologic features are found in the lower dermis or in the subcutaneous fat. If several types of lesions are present and the diagnosis hinges on the histologic findings much time may be saved by taking specimens for biopsy from more than one lesion

In the author's experience a specimen obtained with a 6 mm biopsy punch nearly always has proved adequate for histologic study. Two sutures are sufficient to close the wound

Before placing the specimen in the routine fixative which in many hospitals is Zenkers or Helly's solution one should consider which stains are desirable. Some of the special stains can be performed only if the specimen has been fixed in the appropriate fixative. A tabulation of staining methods with the required fixatives is found in Table 1 on page 29.

LIMITATIONS OF HISTOLOGIC DIAGNOSIS

Mithough histologic study is one of the most valuable means of diagnosis in derinatology, it has its limitations Often no definitive diagnosis can be mide. The reason for this is that few derinatoses aside from the tumots are associated regularly with a diagnosite his

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Myelosis

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Embryology of the Skin

THE EPIDERMIS

In the earliest period of fetal life, the epidermis consists of a single layer of cells. Between the fifth and the seventh weeks of fetal life this becomes a double layer, consisting of an inner layer, the stratum



Fig. 1 The kin of an embry 4 months old. The epiderms consists of three layers the stratum germinatism (\$C) the stratum intermedium (\$1) and the periderm (P). Two primary epithelial germs (£EG) are shown. The fetal derms shows many more fibroblasts it an the adult derms (\$400).

germinativum or palisade layer and an outside layer the periderm or epuriculal layer. The stratum germinativum is composed of large cuboidal cells while the periderm consists of flat cells. In the third month isingle cells appear between the two layers and later form a complete line the stratum intermedium (Fig. 1). The cells of the

3

tologic picture. Instead, the histologic features may be merely sug gestive of a diagnosis or may be entirely nonspecific Even in the case

2

Introduction

of tumors, difficulties in diagnosis may arise For instance, distinction

of squamous cell carcinoma from pseudo epitheliomatous hyperplasia is not always possible. In cases of infectious granulomas such as

syphilis, tuberculosis and the deep mycoses, a specific diagnosis often

cannot be made unless the causative organism can be demonstrated

make a diagnosis possible

cluding a differential diagnosis

and parapsoriasis it is always nonspecific

sometimes it may be merely suggestive. In other diseases of this group, such as the various types of dermatitis or eczema the histologic pic ture is at best, only suggestive. In still others, such as pityrissis roser

Nevertheless frequently, when the histologic picture is not diag nostic, a correlation of the histologic with the clinical findings will

In many instances the chief value of histopathologic study lies in corroborating the clinical diagnosis or in ruling out possible diseases which are being considered on the basis of clinical appearance. It is obvious that the histopathologist can give the clinician a maximum amount of information only if every specimen submitted for histo logic diagnosis is accompanied by detailed clinical information, in

thematosus in which the histologic picture is diagnostic as a rule

Great difficulties may also be encountered in the histologic study of the large group of noninfectious inflammatory dermatoses. In the diseases of this group, such as psoriasis lichen planus and lupus ery

except in a few areas where they persist (See The Apocrine Glands *

page 16)
Melanocytes (Dendritte or Clear Cells) Whereas in the past
the melanin producing dendritic or clear cells were considered to be modified basal cells there is now almost complete agreement that they are neural cells which originate in the neural crest and migrate from there with the nerves to the epidermis during early fetal life (DuShane Rawles) Although this origin of the melanocytes has not been proved experimentally in man it has been proved in mice by Rawles who transplanted tissue from the neural crest of mouse em bryos to the coelom of albano chicken embryos and observed the development of pigmented melanocytes

In human embryos argentaffine melanocytes first are identified in human emotyos argentatine metanocytes tist are identified within the epidermis early in the third month. They possess long brunching dendritic processes which in subsequent months become more numerous and longer up to 100 microns in length joining neighboring melanocytes (Zimmermann and Comblect) (For a further discussion of the melanocytes see pages 9–12)

THE DERMIS

The dermis is of mesodermal origin. During the first months of fetal life it consists of closely packed spindle shaped cells (mesen chymal cells) During the third month fibrils appear at first as a delicate anastomosing argyrophilic network (reticulum fibers). As the fibers increase in number and thickness they arrange themselves in bundles which no longer can be impregnated with silver and which instead begin to stain with the methods for collagenous fibers (Maxi instead organ to stain wan the meanings for consecuous mores quartimon and Bloom). Simultaneously the mesenchymal cells develop into fibroblasts. The elastic fibers appear much later than the collagenous fibers usually in the sixth month (Lynch). The subcuta neous fat first becomes apparent in the third month

It is not as yet settled whether collagen and elastin develop

intracellularly by a direct transformation of living substance of intractinatis by a uncer transformation of tiving substance of mesenchymal cells or extracellularly by precipitation of the ground substance under the influence of an enzyme derived from mesenchymal cells. Observations on the development of fibers in tissue cultures favor the theory of their extracellular origin (Maximow and Bloom BIBLIOGRAPHY

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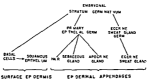
Embryology of the Skin

stratum intermedium are large and because of their clear cytoplasm have a ballooned appearance

At about the fourth month the periderm separates to aid in the formation of the vernix caseosa. At the same time, the stratum intermedium becomes multilisered and develops into the squamous cell layer or stratum malpighii Intercellular bridges become recogniz able only after the epidermis has become stratified with several layers

The embryonal stratum germinativum differentiates into the fol lowing types of cells (1) the basal cells (2) the eccrine sweat gland germ cells and (3) the primary epithelial germ cells (Chart 1)

CHART 1 -EMBRYOLOGY OF THE EPIDERMIS



Basal Cells While the cells of the embryonal stratum germi nativum possess no intercellular bridges the mature basal cells do By progressive differentiation the basal cells develop into squamous cells granular cells and horny cells thus forming the multilayered surface epidermis

Epidermal Appendages The epidermal appendages develop from the primary epithelial germ cells and the eccrine sweat gland germ cells. These cells possess no intercellular bridges and form cells with out intercellular bridges Primary epithelial germs first appear in the third month of fetal life as epithelial buds projecting into the dermis (Fig. 1) Eccrine sweat gland germs are first observed in the fifth or the sixth month

> rnee entolands atrix nd its

two inner sheaths (the Huxley and the Henle layers) develop from the hair matrix. The outer sheath of the hair and the ducts of the sebaceous glands which are composed of prickle cells originate from cells with the potentiality of forming prickle cells namely from basal cells (Lever) The apocrine glands which begin to form with the hair and the sebaceous gland involute before they reach full development

except in a few areas where they persist (See The Apocrine Glands *

page 16)

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It is not as yet settled whether collagen and elastin develop intracellularly by a direct transformation of living substance of mesenchymal cells or extracellularly by precipitation of the ground substance under the influence of an enzyme derived from mesen thymal cells Observations on the development of fibers in tissue cultures favor the theory of their extracellular origin (Maximow and Bloom)

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Histology of the Skin

THE FPIDERMIS

In histologic sections of normal skin, the border between the epidermis and the dermis is irregular because of the fact that numerous cone shaped dermial papilitie reach upward and indent the unier surface of the epidermis. The ridges of epidermis separating the papilite appear in histologic sections as pegs, and, therefore, often are referred to as rete pegs although the term rete ridges is preferable.

Layers of the Epidermis. The epidermis may be divided into four

THE BASAL LAYER Two types of cells occur in the bisal layer

basal cells and melanocytes

Basal cells are columnar in shape and he with their long axis yer

teal to the dividing line between the epidermis and the dermis. They have a deeply bisophilic cytophism and a darkly, straining oral or elongated nucleus. In section, stained with routine istains, the basal cells seem to contain melanin granules often concentrated by

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6 Embryology of the Skin

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Histology of the Skin

THE EPIDERNIS

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Layers of the Epidermis The epidermis may be divided into four

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teraur

located between the granular and the horny layers. The cells in the various layers represent different stages in the gradual evolution of the basal cell into cormified cells and do not represent different types of cells.

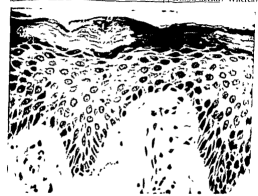
THE BASAI LAYER Two types of cells occur in the bisal layer basal cells and melanocytes

Basal cells are columnar in shape and he with their long rxis vertical to the dividing line between the epiderius and the dermis. They have a deeply broophilic extoplasm and a datkly stanning or all or elongated nucleus. In sections stained routine stains, the break cells seem to contain nelanin gran beautiful.

their nuclei as supranuclear and Montgomery using silve

s a greater number of mitotic figures

epidermis to the dermis is accomplished by the firm interlocking of cytoplasmic processes of the basal cells with reticulum fibers located in the uppermost dermis. Whereas,

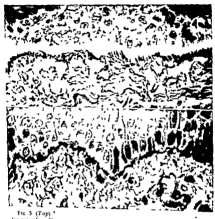


His 2 Normal epidermis, dorsum of the hand Tour layers can be recog nized (1) basil layer (2) stratum malpighii (3) granular layer and (1) horny layer No stratum lucidum is present Note the presence of intercellular bridges between the bisil cells. Several clear cells (melanocytes) are present in the bird layer. They possess a small dark nucleus and clear cytoplasm $(\times 400)$

in perpendicular sections, these reticulum fibers appear like the bristles of a brush (Lig 3), horizontal sections reveal them to form a continuous fibrillar meshwork around the cytoplasmic processes of the basal cells (Odland). In contrast with formerly expressed views, clastic fibers do not contribute to the coherence of the dermis and the epidermis since they do not extend high enough to reach the epi dermis (Diek)

Whereas routine stains do not show a basement membrane, stains with the periodic acid Schiff stain after Hotchkiss and McManus (sec page 30) show a thin homogeneous dense band (Lig 1) at the dermal epidermal junction, indicating the presence of a relatively large amount of polysaccharide material in this zone (Stoughton and

Wells) This band represents a relative barrier to the diffusion of large particles as proved by the fact that dye injected intradermally beneath a subepidermal bulla will not appear in the bulla fluid al



for reticulum fe wirk into which in a firm attache

Fig. 4 (Bottom) Junction between the et ideem can't 1

" (A30)

tho at The ban

Melanocytes (Clear or Dendritte Cells) Melanocytes are of neural origin (see Embryology page 5) They stain with Bloch's dopa

stain (because they have the ability to form melanin) and usually also with silver stains (because they contain melanin) (see page 12) They also strin with gold. In sections strined with hemitoxylin and eosin melanocytes appear as clear cells having a small dark staining nucleus and a clear, slightly basophilic cytoplasm (Figs 2 and 254) On the other hand, in sections impregnated with silver they appear as den drittic cells with numerous long, branching processes provided a suffi cient amount of melanin granules are present within the processes to show their outline (Ebert, Becker Jr, Fitzpatrick and Mont gomers)

THE STRATUM MALPIGHII The cells of the stratum malpighii which are called squamous cells or prickle cells, are polygonal and form a mosuc. They become llattened toward the surface (Fig. 2) The cells are separated by spaces which are traversed by intercellular bridges or prickles. They separate the squamous cells so that lym phrtic fluid can circulate around them and supply them with nour ishment. The intercellular bridges are formed by the cytoplasm of adjoining cells and tonofibrils which extend through them from cell to cell As studies by pliase contrast and polarization microscopy have shown, tonofibrils pass uninterruptedly from one cell to another all the way from the basal cell layer to the stratum corneum Roentgen spectography suggests that the tonofibrils consist of keratin (Nele mans Keuning van Rijssel and Ruiter)

Each of the intercellular bridges in the stratum malpighti contains a small nodular thickening called the nodule of Bizzozero These nodules are stained selectively by Heidenham's iron hematoxylin which does not stain the epidermal cells and bridges (Tavre). This special strining reaction suggests that the nodules of Bizzozero are special structures and not just a thickening of the intercellular bridges Nieuwmeijer on the other hand believes that they merely represent an optical effect produced by the crossing of tonofibrils in the intercellular bridges

If a gold stain is employed one may observe interspersed between the cells of the stratum malpighii gold impregnited cells with den not obsert ad

so far about their nature. Masson regards them as worn out in crimo cytes which are being carried anal passively toward the surface Ferreira Marques points out that schwannian cells generally are auro

represent intra epidermal receptors of pain Becker Ji and Montgomery, on the other hand regard the Langerhans cells as ordinary melanocytes and maintain that they appear to lie in the

superficial epidermis only because of distortion and wrinkling of the tissue produced by the acid used in the gold impregnation. In separated epidermis, which is less distorted by acid than whole skin, they found gold impregnated cells only at the junction of the epidermis and the detims and never in the upper layers of the epidermis.

THE GRANDLAR LANER The cells of the granular layer are dia mond shaped and filled with granules (Fig. 2). The granules are

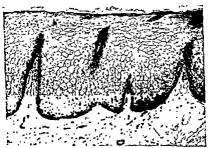


Fig. 5 Epithelium of the oral mucosa. No keratin is formed. The epithelial cells in their migration from the basal layer to the surface first become vacuolated then shrink and finally desquamate. (X200)

coarse, tregular in size and shape, and strongly light refractile. The chemical nature of the granules is not known. Unna called them heratohydme granules, but the substance differs in its chemical behavior from both keratin and hyalin. The thickness of this layer

٩(

mm on the flexor surface of the forearm to 0.5 mm and more on the soles " 1 cells; they are closel 1 in this lan.

The mucous membrane of the mouth normally possesses no gran

ular cells and no horny layer There the epithelial cells in their migration from the basil layer to the surface first become vacuolated, then shrink and finally desquamate (Fig. 5)

THE STRATUM LUCIDUM This layer occurs only on the palms and the soles. It is located above the granular layer and is composed of two or three layers of flat, anucleur cells of homogeneous, transparent appearance. The substance filling the cells and groung them the oily appearance is called cleidin, which is believed to be formed by lique faction of the keratohyaline granules. Ordinarily, this layer cutnot be demonstrated in sections stained by routine straining methods. However, it can be shown by two rather complicated staining methods described by MacLood and Murende.

Pigment of the Epidermis For the study of cut meous pigmenta tion, two stams are necessary the silver stum and the dopt stain Silver stains melanin black, thus indicating its presence Dopt scuns

and~

Becker, Praver and Thatcher, consists in bathing sections of skin in a 0.1 per cent solution of levorotatory 3,4 dihydroxyphenylahinine (called "dopa" for short). Following Bloch's suggestion, the cells that stain dark with the dopa stain are called dopi positive. Such cells stain dark because the enzyme tyrosinase which they contain changes the colorless dopa of the stain into a dark insoluble product, the dopa melanin (Bloch, 1927). The dopa stain closely imitates physio logic melanin formation, which occurs because the amino acid tyrosine in cells containing the enzyme tyrosinase is transformed to dopa and further to melanin by the action of this-enzyme (Lerner and Fitzpatrick).

The dopa stain shows that in the hasal layer only the neural

di ne.

layer of the epidermis but also are found interspersed between the hair mutricells of the hair bulb. When active melanin formation occurs (e.g., after exposure to the sun) the number of dopi positive melanocytes in the bisal layer of the epidermis increases greatly.

The silver stam does not, like the dopn stam, demonstrate the site of formation of melanin but instead demonstrates the presence of melanin Thus, in the also mesodermal phage

matophores stain pos

dermit melanin is seen only in the region of the hasal lyer. However in a deeply pigmented skin as in the Negro, melanin is found also in the upper layers of the epidermis and in phagocytic cells of the dermit Studies by Becker Jr. Fitzpatrick and Monigomery on separated epidermis have revealed that even in deeply pigmented skin melinin is never found in basal cells or squamous cells. It is concained exclusively in melanocytes and their dendritic processes until the stratum granulosum is reached. At this level the dendrites term intended and free melaning granules are seen in the granular layer and the stratum corneum.

pain have been demonstrated extending upwird from the dermis into the lower layers of the stratum malpighir and at the fingers even into the stratum granulosum (Woollard Weddell and Harpmin). With the methylene blue method, these nerve fibers subserving menhate and account of the stratum granulosum (Woollard Weddell and Harpmin).

ture a company on tourh. They are most abundant on the fingers where they may show connections by means of medul lated fibers with Viessner's tactile corpuscles located in the papillar (Woollard).

Formerly two types of cells within the epiderinis were regarded as nerve end cells the Merkel Ramiter tactile cells and the Langer hans cells. The Merkel Ramiter cell is identical with the clear cell and as with it a mele.

THE PPIDERMAL APPENDAGES

The Sweat or Ectrine Clands Sweat glands are present everywhere in the human shin. They are found in greatest abundance on the palms and the soles and in the avillar They are tubular glands in hose secretory cells during the process of secretion do not change in size or shape and do not release any cellular material into the lumin of the gland. Schiefferdecker called the sweat glands ecceitte glands because of his belief that their secretory cells.

Sweat glands are commonsed.

secretory portion hes codec

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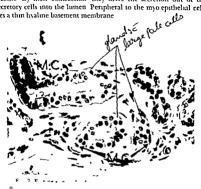
tissue which connects with the subcutaneous fat. Although eccrine glands are formed embryologically with two layers of epithelial cells throughout, only one distinct layer, composed of secretory cells, lines



Fig. 6 Normal skin, back of neck. On the left side of the cllustration, a sweat duct (S D) enters the epidermis. In the center, a large sebaceous gland (S G) leads into a follide enutation of a long of haw. On the right side, a large haw (H) lies within a follicle surrounded by sebrecous glands. An arrector pili muscle (A P) is situated in the obtuse angle of the hair. Beneath the large sebaceous gland, a coiled up eccrine sweat gland (S W G) is present (×50).

the secretory portion in postembryonal life (Fig. 7), because the second, outer layer of epithelial cells has become differentiated into impopping the cells. The secretory cells are large, cylindrical cells with clear, slightly basophilic cytoplasm. The myo epithelial cells are small, spindle-shaped cells inserted sporadically between the secretory cells at their bases. Their long axis extends at a right angle to that of the secretory cells, i.e., parallel with the circumference of the

gland. They possess long processes composed of cytoplasmic fibrils with the same staining properties as smooth muscle and are contractile By their contraction they drive the secretion out of the secretory cells into the lumen Peripheral to the myo epithelial rells hes a thin hyaline hasement membrane



a a protepunctual cells are wedged in at their bases (M.C.) The ducts are composed of two laters of small cuboidal dark staining cells. The lumina of the ducts are lined with a homogeneous cuticle (×400)

The ductal part of the sweat glands leads into the epidermis (Fig 6) Up to the epidermis the duct is composed of two layers of small cuboidal deeply basophilic epithelial cells No peripheral basement membrane is present but the lumen of the duct is lined with a deeply eosmophilic, homogeneous membrane/or cuticle Some authors come dermis ar

cells in

yardy and Memmesheimer, 1936) Others however like Pinkus maintain that sweat ducts have their own lining cells within the epidermis. The observation by Holyoke and Lobitz that the cells lining the intra epidermal portion of the 41

sweat duct maintain their normal structure-when surrounded by carcinoma tends to support Pinkus' contention.

The lumen of sweat glands is small, measuring approximately 90 microns in diameter in the secretory portion, and less still in the ductal portion. In the horny layer of the epidermis, it measures only from 5 to 10 microns in diameter. (The diameter of an erythrocyte is 7.6 microns.)

The Apocrine Glands. The apocrine glands differ from the eccrine plands in origin, distribution, mode of secretion, size and staining reactions. They represent vestignal scent glands



the great difference in the size of the lumina of the apocrine glands (left), and of the eccrine glands (right). The apocrine gland cells show "decapitation secretion (×100)

directly to the epidermis

Apocrine glands are encountered in only a few areas in the axillae, around the nipples, in the perigenital and the perianal regions, and, and field alands in the external ear canal (ceruminous glands), in ie breast (mammary glands) Oc-

found on the abdomen and the

lande

chest.

name apoctine for these glands because part of the cytoplism of their secretory cells is pushed off ()po = off)

because the outer layer has become differentiated just as in the sweat glands into myo epithelial cells. The secretory cells vary greatly in height depending on the stage of their secretory cycle. The secretory cells stam distinctly ensumphilic in contrast with the secretory cells of sweat glands, which stam shiphly beophilic lin addition they frequently contain granules which react posturely to iron string. (Homma) The lumen of the apportune glands is filled with cellular (D)

on the modified skin of the vermilion border of the lips the glans penis the inner surface of the prepuce the labia minora and the they(2)

on of duct

a man. On modified skin, where hair is absent, sebaceous glands lead into follicles devoid of hair. The meibomian glands of the eyelids are

Hair. The hair consists of the hair shaft, composed of keratinized cells, and the hair root, composed of nonkeratinized cells. The hair root terminates in a knoblike expansion, the hair bulb, containing the hair matrix cells. A small connective tissue structure, the papilla,

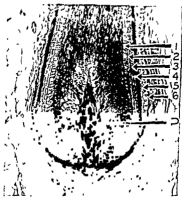


Fig 9 Lower part of a hair. The hair papilla (P) composed of connective tissue, protrudes into the hair bulb The various linings of the hair can be recog nized They are, from the inside to the outside (1) the hair cuticle, (2) the sheath-cuticle, (3) the Huxley layer, (1) the Henle layer (which stains darkly) (5) the outer hair sheath and (6) the vitreous laver (×200)

protrudes into the hair bulb (Fig. 9). The papilla is richly supplied with blood vessels and nerves and, in individuals with dark hair, contains considerable amounts of melanin situated largely in macro phages

Tach hair is surrounded by two inner sheaths and an outer sheath all composed of epithelial cells. The two inner sheaths develop, to gether with the hair, from hair matrix cells, while the outer sheath

represents a downward extension of the epidermis

The hair root consists of a medulla and a cortex. Its cells possess, large, vesicular nuclei and contain variable amounts of melanin Dopa positive melanocytes are interspersed between the hair cells

mis

where the hair rests on the papilla Peripheral to the cortex is the hair caucle which is composed of a single row of nucleated cells (Fig. 9.)

The four shaft is composed of fully keratimized cells closely united with one another and containing either no nuclei or elongited greatly shrunken nuclei. No medulfa and no hair cuttle? The recognizable since they have lost their identity in the process of keratim zation. It may be pointed out that keratimization in the hair proceeds without the interposition of granular cells.

The two inner hair sheaths accompany the hair to approximately the level at which keratinization begins. They are separated from the hair cuticle by the sheath cuticle (Fig. 9.) The sheath cuticle consists of a single layer of c

the cells of the hair cuticle

stain darkly because they ar granules. On this account the

some authors have even stated that the Henle layer contained no nuclei However they can be recognized easily in hair cut on a di agonal plane

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the ep

build he cein not a clear tacuoloted appearance due to the presence of considerable amounts of phones in the extoplasm Outside of the outer hair sheath less a tun membrane of condensed connective trissue the glassy or vireous him (Eg. 9. a)

Nuls. The nari is composed of kerain It grows from the nati matrix which is located beneath the nul fold. The nail matrix con sists of epidermis without a stratum granulosum. Thus just as in the littr ketatimization in the nul proceeds without the interposition of a granular layer. The epithelium of the nul bed has no stratum granulosum either. It is rete rudges are contracted not as a network of anastomosing ridges as elsewhere in the skin, but as parallel long tuthinal nulges.

THE DERVIIS

Three types of fibers occur in the dermis collagenous elastic and Collagenous Education Collagenous Education

Collagenous Fibers Collagen forms more than 98 per cent of the connective tissue of the dermis. It occurs in bundles of fibers held

Hair. The hair consists of the hair shalt, composed of keratinized cells, and the hair root, composed of nonkeratinized cells. The hair root terminates in a knoblike expansion, the hair bulb, containing the hair matrix cells. A small connective issue structure, the papilla.

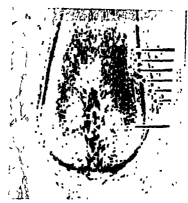


Fig. 9 Lower part of a hair. The hair papilla (P) composed of connective tissue, protrudes into the hair builb. The aurious linings of the hair can be recognized. They are, from the inside to the ousside (1) the hair cuttele, (2) the sheath functe (3) the Huvley layer (4) the Henle layer (which stains darkly) (5) the outer hair sheath and (6) the auricous layer (x200).

protrudes into the hair bulb (Fig. 9). The papilla is richly supplied with blood yessels and nerves and in individuals with dark hair, contains considerable amounts of melanin situated largely in macro phages.

Tach hair is surrounded by two inner sheaths and an outer sheath, all composed of epithelial cells. The two inner sheaths develop, to gether with the hair, from hair matrix cells, while the outer sheath represents a downward extension of the epidermis.

The hair root consists of a medulla and a cortex. Its cells possess, large, vesicular nuclei and contain variable amounts of melanin Dopa positive melanocytes are interspersed between the hair cells

are penetrated more easily by the colloidal silver (Nagcotte and Guyon).

namely, reticulum cells, histiocytes, lymphocytes, vascular endothelial cells, smooth as well as striated muscle cells and fat cells (Dublin).

or sweat grands and around the blood vessels. In addition, retreatum fibers are present in the uppermost dermis (Fig. 3). In perpendicular sections, these retriculum fibers appear arranged like the bristles of a brush, but horizontal sections show a continuous fibrillar mesh work (Odland). Due to the fact that cytoplasmic processes of the

the applicance of reticulum fibers in pathologic conditions in which there is formation of young mesodermal cells. Large amounts of reticulum fibers are present

. . ., occome connetised to collagen

Nerves and Nerve End Organs On sections stained by routine methods, one easily can recognize the large nerves and the Pacini and Messare red organs. The fine nerves and the other nerve end organs require special staining either impregnation with silver salts, e.g. by the Bodium stain (Bodiam), or vital staining with methylene blue (Woollard, Weddell and Harpman)

The skin is endowed with nerves from both the cerebrospinal and the autonomic or vegetative, system. The sensory nerves belong to the cerebrospinal system, while the vasomoro nerves and the nerves supplying the smooth muscles and the unexplained in

gree, since the former up to their terminal ramifications always possess a myelin sheath while the latter usually do not. A few sympa thetic fibers may possess a myelin sheath (Jaeger) together by an interfibrillary ground substance. The fibers and the bundles are correct toward the subcutaneous tissue and finest to ward the outermost portion of the dermis. In the papillary and the subpapillary portions of the dermis, the collagen bundles run in an apparently haphazard manner and do not interface. In the lower dermis, the bundles are arranged nearly parallel to the surface of the skin and do interface. Collagen bundles are only slightly extensible, but, since they are way, they permit stretching of the skin A small number of cells, the fibroblasts, are interspersed between the collagen bundles. They have onal or spindle shaped, pale nuclei en do embrane (see page 36)

bundles are interworen with a web of state shiftly way, and, therefore, only a small portion of any fiber is seen in any histologic section. The fibers are thickest and densest in the lower portion of the derinis, where they are arranged just as the collagen bundles, chiefly parallel to the surface of the skin and are up to 200 microns long. In the uppermost dermis, there are only few small fibers running fortzontally, obliquely and sometimes vertically. From these fibers still finer fibrils spread out toward the epidermis but do not reach it (Dick). Therefore, clastic fibers do not contribute to the attach ment of the epidermis to the dermis, as had been believed formerly.

turn of the skin into its normal position after stretching, thus supplying elasticity to the skin. Their rigidity prevents overdistention When the skin is overstretched as in pregnancy, the elastic fibers may break and degenerate

Reticulum Fibers Reticulum fibers (or lattice fibers "Gitter fasern) form a third system of fibers. They are not visible as such with routine stains but stain with silver (Loots stain). It now is agreed quite generally that.

collagen'), or collagen occu

lagen is merely compacted

easily can see, in sections containing reticulum fibers and stained with Foot's stain, that in areas where the reticulum fibers are densest they tend to aggregate into collagen bundles (Fig. 89). The essential identity of collagen and reticulum is based on the fact that both react alike to all stains with the exception of silver stains (Mallory and Parker). The difference in argy-rophilic properties is due to the fact that reticulum fibrils are finer than collagen fibrils and therefore.

detected easily in microscopic sections (Fig. 10). Located in the subcutis especially of the prlims and the soles they aid in the mediation of the sense of pressure. They are large oval omonike structures composed of a cortex-a core and a myelinated nerve which enters the structure at its lower pole. The cortex consists of from 20 to 60 concentric layers of fibrous tissue the

core consists of semisolid material in which the nerve is embedded. The myelin sheath accompanies the nerve up to the upper pole of the Pacini corpuscle. There the nerve ends in numerous ramifications.

MEISSNER TACTILE BODIES These are located in papillae (Fig 11) and mediate a sense of touch They are present almost exclu sixely on the hands and the feet and are especially numerous on the palmar surfaces of the fingers with their number increasing dis tally On the finger tips they lie in groups of two or three in ad joining papillae thus supplying a spatial relationship to the sense of touch (Weddell) They are cone shaped and he with their long axis perpendicular to the surface of the skin They are of such size that they occupy the greater part of the papilla in which they are located They are composed of a connective tissue capsule and flat



located in a papilla of the fingerup It is composed of flattened (schwan man) cells which have their longest diameter in a transverse direction. Since this is a hematoxylin and coun stain the nerve fibrils cannot be visualized (x400)

tened cells probably schwinnian cells which have their longest drimeter in a transverse direction. One to four impelinated nerves lend to each body. They lose their myclin sheath on entering the body and divide into fibrils which pass upward in a spiral fathion.

RUFFINI BODIES The Ruffini bodies are located in the deeper dermis and in the subcutis Some believe that they mediate a sensation of heat. They present brushlike rumifications of a nonmyelinated nerve within a thirt connective custue capsule.

RRAUSE Bonies. The krause bodies are irregularly shaped spherical formations located in the dermis close to the epidermis. They

22 Histology of the Skin

The basic unit of any nerve is the neurofibril, several of which form a neurite or axon. An axon may or may not have a myelin sheath, but always is surrounded, except at its terminal ramifications, by the neuro-ectodermal schwannian sheath and the mesodermal endoneurium. In medium-sized nerves, several such units are bound together by the permeurium. Large nerve trunks have several such components and an epineurium as a cover. All autonomic nerves and



The 10. Pacini corpuscles in the subcutaneous fat of a fingertip. Their brigginess becomes apparent if one compares their size with that of the ectrine sweat glands and their ducts which are located on the right side of the field (750)

most cerebrospinal nerves end freely in numerous ramifications. However, some of the latter end in special nerve end organs.

Nerve end organs are not present to an equal degree everywhere. They abound in areas of refined sensations, such as the palms and the soles, the lips and the genual region, and are sparse elsewhere. The function of some is not established clearly. Several types of nerve end organs occur the Pacini bodies, the Meissner tactile bodies, the Ruffini bodies, the Kause bodies and the genual corpuscles. These end organs all have a similar structure—a connective-tissue capsule surrounding a core in which the afferent nerve splits up into numerous branches.

PACINI CORPUSCLES, The Pacini corpuscles are the largest of the end organs. They measure up to 2 mm in diameter and thus are

collagen bundles. Lymphatic vessels begin as loops in the papillae and lead down to a lymphatic plexus in the subpapillary layer from which they pass down through the detrmis to a deeper plexus at the junction of the dermis and the subcuits. Lymphatic vessels are lined by only one layer of endothelial cells.

pilorum the tunica dartos of the scrotum and the muscle fibers in the arcola of the nipple are smooth muscles. The muscle fibers of the arrectores pilorum are anchored in the connective fissue of the papillae and are attached to the hair follicles below the sebaceous gland. They are situated in the obtuse angle of the follicle, so, that on contracting they make the follicle more vertical and produce goosefiesb. (Fig. 6)

Strated or voluntary muscle shows cross striation of its filters. The nuclei are located at the periphery of the filters. Strated muscle is found in the skin of the neck (platysma) and the face (muscles of expression)

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have a thin fibrous capsule within which nonmyelinated nerves branch. They are believed to mediate a sensation of cold

GENITAL CORPUSCLES These have the same general structure as the Krause bodies, but are larger and have a thicker capsule

MERKEL-RANVIER TACTILE CELLS These cells, located in the epi dermis, are discussed on page 13

Blood Vessels. The arrangement of the cutaneous blood vessels is inconstant and varies in different parts of the slim. Nevertheless a deep plexus at the junction of the dermis and the subcutaneous tissue and a superficial plexus in the subpapillary layer are always present. The deep plexus has arterioles which possess three layers adventitia, media and intima. The media and the adventitia gradually become thinner as the arteriole ascends through the dermis. The capillaries in the papillary layer are composed merely of one layer of endothelial cells surrounded by a few histocytes (perithelial cells). In some instances, an additional type of cell, called the pericyte, is present. This cell has branching processes which encircle the capillary and are contractile. They can thus change the caliber of the lumen. They represent modified smooth muscle cells (Stout and Murray).

A special structure, the glomus is distributed widely throughout the dermis, but occurs most abundantly on the tips of the fingers and the toes and in the nailbeds. The glomus is concerned with tempera ture regulation and represents a special short circuit device connect ing, without the interposition of capillaries, an arteriole with a venule It consists of an arterial and a venous segment. The arterial segment, called the Sucquet Hoyer canal, branches from an arteriole and has a narrow lumen and a thick wall. The wall consists of an inner endothelial layer and of several layers of glomus cells which are large cells with clear cytoplasm resembling epithelioid cells. The glomus cells are in intimate association with a rich network of non myelinated nerve fibrils demonstrable by silver stain. Although myo fibrils are sparse or absent in the glomus cells they are contractile The glomus cell generally has been regarded as a modified smooth muscle cell (Popoff Weidman) On the basis of tissue culture experi ments, Murray and Stout believe that the glomus cell is derived from the pericyte The venous segment of the glomus has a wide lumen The blood is emptied from the venous segment into sub papillary venules and through the latter into deeper veins

Lymphatic Vessels. Lymphatic fluid circulates in the epidermis around the squamous cells, which are kept apart by intercellular bridges or prickles Similarly, lymphatic fluid circulates between the

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TABLE 1—Survey of Staining Methods Employed at the Massachl setts
General Hospital

FIXATION AND EMBEDDING	STAIN	PURPOSE OF STAIN	RESULTS
Helly s-paraffin (Autotechnicon)	Hematoxylin and cosin t	Routine	Nuclei blue, collagen, nerves, muscle red
Helly s-paraffin (Autorechnicon)	Phosphotungstic acid hematox ylin	Nuclear details and for distinction of collagen and muscle	Aucles, nerves, muscle blue, collagen brown
Helly s-paraffin (Autotechnicon)	An line blue (Mallory)	Collagen	Nucles, nerves, muscle bright red, collagen blue
Helly s-paraffin (Autotechnicon)	Masson s tri chrome	Collagen	Nutlet, nerves, muscle dark ted collagen green
Helly s-paratin (Autotechnicon)	Van Gieson	Collagen	Collagen red, nucles, nerves, rnuscle yellow
Helly s-paraffin (Autotechnicon)	Verhoeff	Elastic tissue	Elastic tissue black
Helly s-paraffin (Autotechnisun)	Foot	Reticulum	Renculum and neurofibrils black
Helly s-parafin (Autotechnicon)	Potassium ferro	lron	Iron blue
Helly s—parafin (Autotechnicon)	Giema	Eounophils, bacteria, Donovan bodies, Lesshmansa, Frisch bacilli, ffistoplasmi capsulatum	sinophils, Donovan bodies,
Helly s-paraffin (Autotechnicon	Gram,	Bacteria	Gram positive bacteria blue, Gram-negative bacteria red
Helly s-paraffin (Autotechnicon	Ziehl Seelsen	Acid fast bacteria	Acid fast bacteria red Acid fast bacteria black
10° Formalis— paraffin	Levaditi	Spirochetes	Sparochetes black
10° c Formalis — paraffin	Bodisn	verves	Neurofibrils black
10° o Formalin —	N. L 1 L	.11	· · ·
10° Formalin- frozen section	-		Chalesterol brownish red
10° Formal n- frozen section	Crystal violet	Amyloid	Ams loud purple
Absolute alcohol ceiloidin	Muticarmine	Mucia	Mucan red
80°c alcohol- paraffin	Von Kossa	Calcium	Calcium black
Fresh tissue frozen section	Dopa	Melanin producing cells	Melanin producing cells show dark granules

4

Laboratory Methods

FIXATION, EMBEDDING AND STAINING

Fixation. Helly Mixative is used routinely in the Pathology Laboratory of the Massichusetts General Hospital For its preparation two solutions are required (1) Zenker's solution

> Potassum dichromate 25 Gm Mercury bichloride 80 Gm Water 1000 cc

and (2) Formalin (which is a 40 per cent aqueous solution of formal dehyde) Formalin Lec. is added to each 20 cc. of Zenker's solution just before use. The specimen should remain not less than 5 hours and not longer than 21 hours in Helly's fixative. It is therewashed in running water for at least 2 hours and wansferred into 80 per cent alcohol, in which it can be left indefinitely.

For specimens which have to be trailed, the fixative of choice is 10 per cent Formula because specimens may remain in this fixative in definitely. One cc. of Formula is added to each 9 cc. of water just

before use

Pielly's fixative is contraindicated if one desires to demonstrate lipids, nerve fibers, amyloid, granules of mast cells, spirochetesocal cium origination, or of one vishes to perform aghopa strin. For the demonstration of pids, herve fibers, amyloid, granules of mast cells and spirochetes, fixation in 10 per cent Commign is required, to demonstrate alcum, 80 per cent alcohol is used, and for demonstration of plucin absolute alcohol is the approved fixative. For the fiops state, not fixative should be employed the fresh tissue should merely be wrapped in most gaute and sent to the laboratory.

Dehydrating and Embedding. Following fivation, all routine specimens are placed successively into several beakers containing diovane in increasing concentrations for dehydration and several beakers of hot paraffin for embedding. This may be done by hand or with the aid of an Autotechnicon machine.

In the Pathology Laboratory of the Massachusetts General Hos pital as in most other such laboratories all routine specimens,

TABLE 1—Survey of Staining Methods Employed at the Massachusetts General Hospital

FIXATION AND EMBEDDING	STAIN	PURPOSE OF STAIN	RESULTS
Helly s-paraffin (Autotechn con)	Hematoxyl n and eosin	Routine	Sucles blue collagen nerves muscle red
Helly s-paraffin (Autorechn con)	Phosphotungstic ac d hematox 3 i st	Suclear deta is and] for d st netion of collagen and muscle	Nucles nerves muscle blue collagen brown
Helly s-paraffin (Autotechn con)	An) ne blue (Mallory)	Collagen	Nucle nerves muscle bright red collagen blue
Helly s-paraffin (Autotechn con)	Ma, son s tri chrome	Collagen	red collagen green
Helly s-paraffin (Autotechnicon)	Van C eson	Collagen	Collagen red nuclei nerves muscle yellow
Helly s-paration (Autorechn con)	1 erhoeff	Elastic tissue	Elastic tissue black
Helly s-paraffin (Autotechn con)	Foot	Ret culum	Reticulum and neurofibrils black
FI	p	7	·

		Le shman a Frisch bacill Histoplasma capsula um	s sup Do ovan bod es L2 shman a Frisch barills H stoplasma capsulatum brill ant sed
Helly's paraffin (Autotechnicon)	Gram	Bactens	Gram positive bacteria blue Gram negative bacteria red
Helly s—paraffin (Au o echn con)	Ziehl Neelsen} Fite	Ac d fast bacteria	Acid fast bacteria red Acid fast bacteria black
10° c Formal n- paraffin	Levad tı	Sp rochetes	Sperochetes black
10° o Formal n paraffin	Bod an	*erves	Neurofibrils black
10° o Formal n paraffin	Methylene blue }	Granules of mast cells	Mast cell granules purple
frozen section	Scarlet red	Lipids	Neutral fat bright orange tholesterol brown sh red
10° Formal n- frozen section	Crystal violet	Amylo d	Amylo d purple
Absolute alcohol cello d n	Mucicarm ne	Mucin	Mucin red
80° alcohol paraffin	Von Kossa	Calc um	Calcium black
Fresh tissue - frozen section	Dops	Melan a producing cells	Melan n producing cells show dark granules

whether unfixed or already placed in Helly's fixative or in 10 per cent Formalin, are sent through the Autotechnicon. This machine, which is controlled by an electric clock, accomplishes fixation, delay dration and embedding automatically overnight. The specimens are placed in small, perforated cassettes, which are then put into a per forated metal basket suspended from an arm of the machine. The metal basket is automatically lowered into and raised out of a succession of beakers containing Helly's fixative, water, 95 per cent alcohol, dioxane and warm liquid paraffin.

Specimens that are to be stained for lipids, nerve fibers, amyloid calcium, mucin, spirochetes or granules of mast cells, or are to be treated with the dopa stain, are not sent through the Autotechnicon, in order to/avoid exposure to Helly's fixative Specimens to be stained for herve fibers calcium, granules of mast cells or spirochetes, after fixation, are carried by hand through dioxane and embedded in paraffin Specimens to be stuned for mucin, after fixation are embedded in celloidin Specimens to be stained for fipids or imploid, after fixation, are cut on the freezing nucrotome and then stained Specimens to be stained with dopa are cut on the freezing microtome without previous fixation

Staining All routine specimens, after hiving been cut on a rotary microtome into sections from 5 to 8 microns thick, are stained with hematoxylin and eosin. With this method all nuclei stain blue, and collagen, muscle and nerves stain red. Special stains are employed for the demonstration of particular structures (see Table 1).

HISTOCHEMICAL STAINING

Brief mention may be made of two histochemical stains which have attained considerable practical importance the Feulgen reaction and the Hotchkiss McManus stain (DeLamater, Mescon and Barger) Both stains can be carried out on material fixed in 10 per cent Formalin

The Feulgen reaction results in red staining of desoxyribonucleic acid (DNA) which is present in nuclei and in many virus inclusion bodies. On the other hand, ribonucleic acid (RNA), present in nu cleoli, cytoplasm and keratohyalin, does not stain. This reaction is important as a stain for viruses since it often allows their differentiation from nucleoli and keratohyaline granules.

The Hotchkiss-McManus stain, or periodic acid Schiff (PAS) strin, demonstrates the presence of polysrccharides such as glycogen, hyal uronic acid and other mucoid substances by staining them red. It is also of value in the study of fibrinoid degeneration where there is depolymerization of collagen resulting in the formation of polysaccharides. For the demonstration of glycogen, it is necessary to

compare two serial sections one exposed to hastase prior to staining and the other not. Since the split products of glycogen resulting from the action of the disastase no longer are colored red by this stain only such substances represent glycogen which stain red with out preliminary disastase digestion but do not stain after disastase digestion.

In addition the Hotchkiss McManus stam is of great value in the demonstration of fung. Since the cell walls of fung are composed of a mixture of cellulose and churn they contain considerable amounts of polysaccharides. All fungi therefore stain bright red and thus are detected easily in histologic sections because other tissues stain very faintly or not at all (Aligman Mescon and DeLamater).

POLARISCOPIC EXAMINATION

Polariscopic examination is the examination of histologic sections under the microscope with polarized light—i.e. with light from which all rays except those vibrating in one plane are excluded

For the polariscopic evamination two disks mide of polarizing plastics are inserted into the microscope. One disk is placed below the condenser of the microscope und acts as the polarizer. The second disk is placed into the eyepiece of the microscope and acts as the analyzer when the eyepiece continuing the analyzing disk is rotated so that the path of the light through the two disks is thorated at a right angle the field is dark. When however doubly refractile lipids are introduced between the two disks they break the polarization and are visible as bright white bodies in the dark field.

Cholesterol and cholesterol esters are doubly refractule while neutral fats are not Doubly refractile fats are called anisotropic the others isotropic Doubly refractile lipids are present regularly in the cutaneous lesions of xanthomatosis and hyperlipening in xanthelasma palpebrarum and in extracellular cholesterois. They are present occasionally in Hand Schuller Christian disease in foreign body granulomas and in histocytoma. The lipid material in lipid pro teinois and necrobiosis lipidica diabeticorum however is not doubly refractile as a rule.

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5

Morphology of the Mesodermal Cells

Various types of mesodermal cells infiltrate the dermis and occasionally the epidermis in the inflammatory dermatoses, in the granu lomas and in the lymphomas It is important for diagnostic purposes to identify the cell types. Three groups of mesodermal cells are recognized generally the myeloid group, the lymphoid group and the reticular (or histocytic) group. In addition, plasma cells and mast cells occur. It is likely that both originate from the reticular group of cells.

MYELOID GROUP

Polymorphonuclear leukocytes and eosinophilic leukocytes may occur in the skin in various dermatoses. Basophilic leukocytes how ever, do not occur. In myelosis (myeloid leukemia), in addition to polymorphonuclear leukocytes and eosinophilic leukocytes one finds immature myeloid cells—namely, myeloblasts and myelocytes.

Myeloblast and Myelocyte. The myeloblast is a large cell with non granular cytoplasm and a round or oxal, resicular nucleus. The myelocyte differs from the myeloblast mainly by having in its cyto plasm either neutrophilic or eosinophilic granules. Myelocytes are always oxidase positive, while myeloblasts may be oxidase positive or oxidase negative. (For a more detailed discussion, see page 191)

Polymorphonuclear Leukocyte The polymorphonuclear leukocyte, or neutrophil, has a lobated nucleus and contains fine neutro philic granules in its cytoplasm This cell occasionally is referred to as a microphage because it is a small phagocytic cell with the ability to phagocytice becteria only In contrast, the macrophage, a large phagocytic, histiocytic cell, also can take up particulate matter such as hemosiderin, melanin and lipid Because of their ability to phago cytize bacteria, mimerous polymorphonuclear leukocytes are present in the skin in acute bacterial infections, e.g., in erysipelis and folliculities. In addition, they are found in conspicuous numbers in acute dermittits, allergic vasculitis (anaphylactoid purpura), crythema

erythema nodosum and

illed polymorphonuclear

cosmophil is characterized by the presence of coarse round bril liant cosmophilic granules in its cytoplasm. The granules are visible with rounne stains but stand out more clearly when Gremya's stain is used. Its nucleus is lobated and thus has the same appearance as that of the polymorphonuclear leukocyte. The cell is able to phagocytic bacteria. Polymorphonuclear eosinophils form as such in the hone marrow.

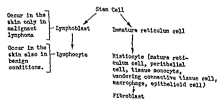
In addition to polymorphomiclear eosinophils monolobed eosinophils are observed not infrequently in pathologic conditions of the skin (Burkhart and Montgomery). Their nuclei are either oxal kidney shaped or band shaped. It is possible that they represent his stocytic eosinophils and as such are formed in the skin.

Both the polymorphonuclear and the monolobed cosmophil occur in the tissue as a response to local anaphylaxis (Campbell) Tissue cosmophila is apit to be prominent in eczematous drug eruptions atopic dermatitis dermatitis herpetiformis pemphigus vegetans alore dermatitis (anaphylactoid purpura) ecomophilic granuloma granuloma faciale mycosis fungoides and Hodgkin is disease

LYMPHOID GROUP

It is widely accepted that lymphoid and reticular cells arise from a common cell—the lymphoid reticular stem cell (Chart 2)

CHART 2-DEVELOPMENT OF LYMPHOID AND RETIGULAR CELLS



Stem Cell. The stem cell can be seen in the ?

matin particles and one or two prominent nucleoli (see page 471 and Figure 260). This cell may develop into either a lymphoid cell (stem cell \rightarrow lymphoblast \rightarrow lymphocyte) or a reticular cell (stem cell \rightarrow reticulum cell \rightarrow histocyte \rightarrow fibroblast) (Gall and Mallory)

Lymphoblast. Lymphoblasts do not occur in ordinary inflammation of the skin but only in lymphoma. They are spherical cells and possess a large nucleus surrounded by a uniform, bisophilic rim of cytoplasm. The nucleus is round or slightly indented. (see page 477 and Figure 263). It is much larger and lighter than that of the mature lymphocyte.

Lymphocyte. Lymphocytes have much smaller nuclei and less cyto plasm than lymphoblasis. The nuclei are spherical and appear very dirkly stained since the chromatin forms a thick layer at the periph ery and several dark staining particles in the interior.

Lymphocytes are found in the skin in most forms of lymphomi and in actite, subacute and chronic inflammations—for instance, in acute, subacute and chronic dermatitis, in psoriasis in lichen planus and in lupus erythematosus. They also form a large proportion of the cells in most granulomas.

It is probable that the lymphocytes present in the skin are hematog enous in origin, except in malignant lymphoma, in which they may arise autochthonously. It is assumed by some authors that lymphocytes may transform into histocytes (Kolouch). Some believe even that they may transform into fibroblasts (Goldsmith).

RETICULAR OR HISTIOCYTIC GROUP

The reticular or histocytic group of cells belongs to the reticulo endothelial system. The two outstanding properties of the reticular or histocytic group of cells are their ability to absorb particulate matter and to produce reticulum fibers. As the cells age they may change into fibroblasts (see Chart 2, page 33). As such, they no longer possess phagocytic powers and form collagen rather than reticulum.

No generally accepted nomenclature exists for the reticular group of cells. The following names are used by various authors as designation for the mature cell of this group. Instituocyte, reticultum cell, clasmatocyte, perithelial cell, tissue monocyte and wandering connective tissue cell. The term histocyte at present is employed the most widely. Under special conditions, the histocyte may become a macrophage or an epithelioid cell.

Immature cells of the reticular groups are observed in lymphoma and generally are referred to as reticulum cells. Therefore, lymphomas in which these cells predominate are called reticulum cell.

lymphoma However, it should be kept in mind that many authors use the term rejiculum cell also for mature rejicular cells

Reticulum Cell (Immature Reticular Cell) This cell, as seen in reticulum-cell lymphoma possesses a nucleus which is smaller than that of a stem cell but larger than that of a histocyte. The nucleus may be round but usually is oval or kidney shaped. It is pale strining and has a moderately heavy chromatin network and a distinct basephilic nuclear membrane (see page 174 and Figure 261) The cyto-plasm is eosinophilic and abundant. It often is irregular in outline and may show pseudopodic protrusions. Because of their immaturity. the cells may form reticulum fibers only to a slight degree and have little or no phagocytic power

Histocyte (Mature Reticular Cell) Histocytes are formed in the skin in contrast to lymphocytes which do not arise in the skin except in malignant lymphoma

In the normal skin histocytes are present in small numbers around the capillaries (perithelal cells) Under pathologic conditions which include the inflammatory diseases and the granulomas histiocytes tend to wander from the pericapillary regions into the dermis (tissue monocytes wandering connective tissue cells)

Histocytes are much smaller than reticulum cells but otherwise

do not differ from them in appearance Histocytes form abundant reticulum fibers and possess the ability to phagocytize bacteria and particulate mairer

Histocytes may resemble fibroblasts greatly and often it is not possible to state with certainty whether a given cell is a histocyte or a fibroblast. As a rule however the nuclei of histocytes are larger and stain slightly paler than those of fibroblasts. Although both may have oval nuclei those of histocytes tend to be kidney shaped those of fibroblasts spindle shaped. While histocytes possess the ability to form reticulum fibers fibroblasts form collagen. It should be remem bered that no sharp borderline exists between the histocyte and the fibroblast since the histocyte presents the parent cell of the fibroblast and may develop into a fibroblast

Macrophage Histocytes possess the ability to phagocytize particu stationage transfer poses the agint to pragocytic particulate matter and certain micro organisms. Those containing phagocy tized material are called macrophages. Histocytes migrate as wan dering cells to are is where material digestible to them is present or melanophor

They may also

bacteria fungi may be ingested by histocytes are lepta bacilli and Frisch bacilli of

fungi, Histoplasma capsulatum; of viruses, Donovan bodies, and of protozoa, Leishmania tropica When individual macrophages are unable to deal with particles to be removed, they tend to finse io gether and to form multinucleated foreign body giant cells Fycel lent examples of foreign body giant cells may be seen in paraffinoma gout and Malherbe's calcifying epithelioma. The nuclei in foreign body giant cells usually are clumped together in an irregular arrangement, but they may he regularly along the periphery of the cell, so that foreign body giant cells may be indistinguishable from Langhans giant cells

Epithelioid Cell. Under certain conditions, histocytes may change into epithelioid cells. Epithelioid cells possess a large, usually oxal, pale, vesicular nucleus resembling the nucleus of epithelial cells and abundant, ill defined, slightly eosinophilic cytoplasm. Pseudopodic elongations of the cytoplasm usually can be seen. When lying in groups, the cytoplasm of neighboring cells often appears collesced Epithelioid cells may form giant cells, the so called Langhans type of giant cell. It is likely that this type of giant cell forms by amitous nuclear division without cellular division. The nuclei are arranged in an arc along the periphery of the cell in horseshoe fashion. Epithelioid cells and Langhans giant cells have phagocytic power and have the ability to form reticulum fibers.

Epithelioid cells are found in a variety of granulomas, especially in tuberculosis, sarcoidosis, leprosy, spihlis blistomycosis and cocci dioidomycosis. In tuberculosis epithelioid cells form as a tissue re-sponse to the lipid fraction of the tubercle braillus (Sibin). As a granulomatous lesion containing epithelioid cells heals, the epithelioid cells may mature into fibroblasts and their reticulum fibers into collagenous fibers. This process can be observed particularly well in healing lesions of sarcoidosis.

Fibroblast. The common (resting) connective tissue cells are called fibroblasts because they are instrumental in the elaboration of the collagenous fibers (see page 20). Their nuclei are elongated, often spindle shaped. Because of their pale staining and the presence of a fine nuclear membrane, the nuclei have a vesicular appearance. The cell body, which is spindle shaped is not discerned easily. Extraoblasts usually are found adjacent to the surface of collagenous bundles.

PLASMA CELLS

The plasma cell has abundant cytoplasm which is deeply basophilic, homogeneous and sharply defined. The nucleus is eccentrically placed and round, and along its membrane shows coarse, dark stain.

lished All stages of transition between reticulum cells and plasma cells can be observed in the spleen of rabbus after repeated intravenous injections of horse serum Antibodies are formed in great quantities in reticulum cells during their development into plasma cells while mature plasma cells have already passed the stage of greatest activity (Fagraeus). It is assumed by some that not only reticulum cells but also all multipotent cells of the connective tissue as well as 1 jumphocytes are capable of transforming into a plasma cell (Campbell and Good).

Plasma cells occur in small numbers in most chronic inflammator, diseases of the skin and in larger number in granulomis. They are particularly conspicuous in sphilits granuloma inquitable and thino seleroma. In the presence of many plasma cells, but especially in thinoseleroma round hislane acidophilic bodies so-called Russell bodies may be found within and outside of plasma cells. They form within plasma cells as a phenomenon of degeneration and finally are expelled (Pearse). They may attain a size twice that of a normal plasma cell. (See also page 202.)

MAST CELLS

Mast cells and basophilic leukocytes have nothing in common aside from an identical basophilic metachromatic staining reaction of their granules. The

cyte a myeloid c

and have an ova

and have a lobated nucleus

The grunules of mast cells are soluble in ordinary fixatives such as Helly's and Zenker's fluids and do not stain with hematocylin ces in Ten per cent Formalin absolute alcohol, and saturated basic-lead acetate are suitable as fixatives and methylene blue or Giemsa's stain is suitable for staining the granules. The granules are basophilic (i.e. they stain with basic annihine dyes) and metachromatic (i.e. they may stain in a color different from that possessed by the die). In this sense the blush dye thounne stains the mast-cell granules a reddish violet toluidine blue stains them a purplish red and polychrome metilylene blue stains them ad (Michels). Must cells have two known functions secretion of hjaluronic acid

and secretion of heparin. That hyaluronic acid is present in the mast cell granules is proved by the observations that mast cell granules.

ules strin exactly like hyaluronic acid and no longer show meta chromatic staining after the tissue containing them has been sub mitted to the action of hydrironidase (Asboe Hansen) Evidence for the presence of heparin in the mast cell granules is the similarity of teaction of the two substances to metachromatic staining and the parallelism existing between the amount of extractable henrin and the most cell content of certain organs (Jorpes)

Mast cells occur everywhere in the connective tissue of the body particularly in the vicinity of capillaries and in the walls of larger blood vessels. The normal skin contains relatively few mast cells which are small and spindle shaped. As a rule, they are arranged in groups around the blood vessels but they occur also around the hair follicles and in the papillary layer of the dermis. Their number is increased in many different conditions (Asboe Hansen) For instance the granulation tissue in healing wounds contains more most cells than normally are present. In most atching dermatoses, such as atomic eczema contact derinatitis and lichen planus an increased number of mast cells is present around the capillaries. In lupus erythema tosus in which there is an increase in the amount of hydronic acid in the dermis the number of most cells closely parallels the intensity of metachromatic staining. Also neurolibromas and the stroma of carcinomas of the skin contain numerous mast cells. Thus an in crease in mast cells is of no diagnostic significance, except in urticaria pigmentosa where they occur in tumor like aggregates especially in the upper dermis

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Congenital Diseases (Genodermatoses)

ICHTHYOSIS

Two forms of ichthyosis occur ichthyosis vulgaris and ichthyosis congenita. In ichthyosis vulgaris, the skin is dry and rough and shows



Fig. 12 Ichthyosis vulgaris. There are hyperkeratosis and absence of the granular layer. The stratum malpiglin is thin, the rete ridges in regular. Vlarge keratotic plug is located within a hair follicle. (X100)

scaling, often in the form of large lamellae resembling fish scales Involvement is severest on the extensor surfaces of the extremities, while the flexures usually are spired. Follicular hyperkeratoses are frequently present.

Ichthyosis congenita is a more severe form of ichthyosis than

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ichthyosis vulgaris. The skin is represented by a thick horny cuirass with development of deep fissures.

Ichthyous hystrix is not related to ichthyous but represents an extensive or systematized nevus verrucosus (see page 321)

Histopathology The characteristic lesson of ichthyosis vulgatis is hyperkeratosis with diminution or even complete absence of the granular layer (Fig. 12). This represents an exception to the rule that an increase in thickness of the horny layer is accompanied by an increase in thickness also of the granular layer. The probable reason for this exceptional finding is that in ichthyosis vulgaris there is an inadequate shedding ruther than an overproduction of horny cells. The hyperkeratosis in ichthyosis vulgaris is moderate and not associated with parakeratosis. The stratum malpighii is thinner than normal and the rete indges are dimunished in number some are atropluc while others are slender elongated and branching (doce tailed). The hyperkeratosis usually affects also the upper portion of the hair follicles resulting in large follicular keratotic plugs. The pressure of the keratotic plugs causes atrophy of the lower portion of the follicles and of the sebacrous glands.

Ichthyosis congenita differs from ichthyosis vulgaris only by showing a much greater degree of hyperkeratosis. A stratum granulosum is usually present, but it is not prominent.

CONGENITAL ICHTHYOSIFORM ERYTHRODERMA

Like ichthyosis this disease is characterized by dryness roughness and scaling of the skin. In addition generalized crythema is present and in many cases bullae occur. In contrast to ichthyosis the flex ural surfaces are involved most intensely.

Histopathology In the erythematous areas of the skin one observes pronounced hyperkeratosis with occasional islands of parakeratosis A stratum granulosum is present. Although it varies in thickness it is for the most part hypertrophic. The stratum mahpighin shows acanthosis with irregular elongation of the rete ridges. The upper dermits shows a chronic inflammatory infiltrate.

Examination of bullous lesions has revealed in some cases merely presence of a nonspecific subcorneal bulla (Mackee and Rosen). In others a pronounced ballooning of epidermal cells was noted either in the midportion of the epidermis (Ellis) or in the granular layer (Barker and Sachs). In the latter case the changes in the granular layer resembled those seen in epidermodysplasia vertructiorinis. Ellis believes that the ballooning of epidermal cells is caused by a per sistence of embryonal cells inasmuch as the cells of the embryonal strium intermedium have a ballooned appearance (see page 4)

Differential Diagnosis. Congenital ichthyosiform erythroderma differs from ichthyosis by the presence of a usually hypertrophic granular layer, the presence of acanthosis and the presence of an inflammatory infiltrate in the dermis. In ichthyosis, the granular layer is diminished in thickness or absent, the stratum malpighn is thinner than normal and no infiltrate is present in the dermis (Lay mon and Murphy)

KERATOSIS PALMARIS ET PLANTARIS

This condition, which often is hereditary, is characterized by dif fuse thickening of the horny layer of the palms and the soles Because of the formation of fissures, the condition causes great discomfort to the patient

Histopathology. The histologic picture is nonspecific, showing considerable hyperkeratosis as the only constant change. Occasionally there also are acanthosis and a chronic inflammatory infiltrate in the upper dermis

KERATOSIS PUNCTATA PALMARIS ET PLANTARIS

Multiple, discrete, yellow to dark brown, firm, slightly elevated, conical keratotic plugs, 1 to 3 mm in diameter, are distributed sym metrically over the palms and the soles

Histopathology. One observes a circumscribed hypertrophy of the stratum corneum consisting of a cone shaped plug which invaginates the subjecent structures. Beneath this plug the stratum malpighia is thinned. The dermis is free of any inflammatory infiltrate (Scott, Costello and Simuango)

PACHYONYCHIA CONGENITA

This disease shows dystrophic changes of the nails, palmar and plantar hyperkeratosis, follicular keratosis and leukoplakia of the oral mucosa

A variety of puchyonychia congenita is dyskeratosis congenita. It shares with pachyonychia congenita the dystrophic changes of the nails and the presence of leukoplakia However, there are no hyper keratotic changes of the skin Instead the skin shows atrophy and patches of hyperpigmentation

Histopathology. In pachyonychia congenita one observes hyper keratosis with areas of parakeratosis. There is follicular plugging Occasionally, horn plugs are present also in the sweat ducts (An dreus) The granular layer is hypertrophic. The stratum malpighit shows acouthosis with elongation of the rete ridges. A mild chronic inflammatory infiltrate is present in the upper dermis Dyskeratotic

changes, similar to the corps tonds of Darier's discrse, have been observed in the stratum malpighii of some cases (Andrews, Wright and Guequierre)

Dyskeratosis congenita shows thinning of the epidermis and almost complete absence of rete ridges. In areas in which the skin shows hyperpigmentation, the amount of melanin in the bisil layer is increased and melanophores are present in the upper dermis (Cole, Rauschkolb and Toomey)

POROKERATOSIS MIBELLI

One or several lesions may be present. They show an attophic center surrounded by a raised keratotic wall. The wall has on its top a groove filled with keratotic material.



wall It shows the deep grows which forms the center of the keratotic wall. He groove is filled with a large horn plug the cornoid famella. The cornoid famella has a column of parakeratotic cells in its center. On the right is a normal west door,

Histopathology. The atrophic center shows slight hyperkeratosis, atrophy of the stratum malpighii and fibrosis of the upper dermis. The keratotic wall shows considerable hyperkeratosis and acauthosis. In its center, the keratotic wall presents a deep groove filled by a large horn plug (the 'cornoid lamella'). This cornoid lamella has

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in its center a column of parakeratotic cells. Beneath the parakeratotic column the granular layer is absent elsewhere it is well developed. The dermis underlying the cornoid lamella shows a chronic inflammatory infiltrate (Tig. 13).

Owing to the presence of the cornoid lamelly the histologic picture of porokeratosis is diagnostic. The name porokeratosis is a misnomer since the cornoid lamelly is not necessarily located in the

opening of a sweat duct

VERODERMA PIGMENTOSIM

This disorder is associated with hypersensitivity to ultraviolet light. The lesions occur chiefly on exposed areas of the skin. An early and a late stage occur. In the early stage one observes slight diffuse erythem, with scaling and small areas of hyperpigmentation resembling freekles. In the late stage atrophy of the skin mottled pigmentation and telanguectises are present giving the skin an appear ance similar to that of chronic radiodermatics. Warty growths appear within the atrophic skin, which may develop into carcinoma.

Histopathology. In the early stage the histologic appearance is not always characteristic the diagnosis often can be made from the combination of the histologic changes however. There are (1) hyper keratosis (2) thinning of the stratum malpinghi with atrophy of some of the rete ridges and irregular proliferation and prolongation of others (3) edema of the upper dermis (4) a chronic inflammatory predominantly periviscular infiltrate in the upper dermis and (5) spotted melanin pigmentation of the basal layer with melanophores in the upper dermis.

In the late stage the epidermis shows atrophy in some areas and acanthosis in others. Atypical and multinucleated cells may be seen in the epidermis. The hyperkeratosis and hyperpigmentation already present in the early stage are more pronounced. The upper dermis shows degenerative changes of the collagen and of the elastic fibers of the same type as is seen in senile degeneration of the skin. Thus one observes basophilic degeneration of the collagen and senile elastosis (see page 157). In some areas, the epidermis may show atypical downward growth so that the histologic picture in such areas is identical with that of senile keratosis.

Ultimately squimous-cell carcinomis and occasionally bisal cell epitheliomis and sarcomas develop in some of the lesions

CONGENITAL ECTODERMAL DEFECT

This condition represents an incomplete development of the epi dermis and its appendages. The skin is smooth and glossy. Hair growth is sparse or completely absent. The facies is typical, it shows depressed masal root and bridge, promitient frontal bosses and thick lips. In addition, there may be dental aplasts and dystrophy of the mails. Because of the diminution or the absence of sweat glands, the patient is unable to sweat adequately and therefore is intolerant to heat

Histopathology. There is either a total absence of sweat glands or the presence of rudiments of monfunctioning glands and dutis There is usually a similar deficiency of pilosebaccous sirutures. The epiderius is thinner than normal and there may be hisewise a reduction in the width of the derms. The collagen, the elastic fibers and the blood vessels are normal in appearance, as a rule Examination of the axillary skin in two cases revealed total absence of eccrine glands, while the apocrine glands were developed normally (Sunderman).

ROTHMUND'S SYNDROME, WERNER'S SYNDROME (PROGERIA OF ADULTS) AND PROGERIA OF CHILDREN

These are three different, though related, genodermatoses in which

infancy with

skin shows erythema, scaling, telanguectases and brownish pigmentation so that the appearance of the skin resembles that of poikilo-

undergo atrophy leading on the legs to incerations. Cataracts develop early in adult life

Progeria of children is not familial. It starts several months after birth. The patient develops into a dwarf with a large skill and bird like features. The skin appears alrophic and wrinkled.

Histopathology. No characteristic histologic leatures are seen in the skin in any of the three diseases In all of them, the skin merely shows atrophy of the epiderius, thinning of the collagen bundles in the dermis and atrophy or even disappearance of the cutaneous appendages (Reed). The areas of erythema in Rothmund's syndrome, in spite of their clinical resemblance to poskiloderma atrophicans vasculare, show no inflammatory infiltrate but merely atrophy and telanguerius (Thanhauser).

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in its center a column of parakeratoric cells. Beneath the parakera totic column the granular layer is absent elsewhere it is well de veloped. The dermis underlying the cornoid lamella shows a chronic inflammatory infiltrate (Fig. 13).

Owing to the presence of the cornoid lamella the histologic picture of porokeritosis is diagnostic. The name porokeritosis is a misnomer since the cornoid lamella is not necessarily located in the opening of a sweat duct.

VLKODLEMA BIGMENTOSUM

This disorder is associated with hypersensitivity to ultraviolet light. The lesions occur chiefly on exposed areas of the skin. An early and a late stage occur. In the early stage one observes slight diffuse erythem, with scaling and small areas of hyperpigmentation resembling freekles. In the late stage alrophy of the skin mottled pigmentation and telangiecties are present, giving the skin an appear ance similar to that of chronic radiodermatitis. Warty growths appear within the atrophic skin, which may develop into carcinoma.

Histopathology In the early singe the histologic appearance is not always characteristic the diagnosis often can be made from the combination of the histologic changes however. There are (I) hyper keritosis (2) thurning of the stratum malpighti with atrophy of some of the rete ridges and arregular proliferation and prolongation of others (3) edema of the upper dermis (4) a chronic inflammatory predominantly perivascular infiltrate in the upper dermis and (5) spotted melanin pigmentation of the basal layer with melanophores in the upper dermis.

In the late stage the epidermis shows atrophy in some areas and acanthosis in others. Atypical and multinucleated cells may be seen in the epidermis. The hyperkeratosis and hyperpigmentation already present in the early stage are more pronounced. The upper dermis shows degenerative changes of the collagen and of the elastic fibers of the same type as is seen in senile degeneration of the skin. Thus one observes bisophilic degeneration of the collagen and senile elastosis (see page 157). In some areas the epidermis may show atypical downward growth so that the histologic picture in such areas is identical with that of senile kerticosis.

Ultimately squamous cell carcinomas and occasionally basal cell enitheliomas and sarcomas develop in some of the lesions

CONGENITAL ECTODERMAI DEFECT

This condition represents an incomplete development of the epi dermis and its appendages The skin is smooth and glossy Hair scarring results. In hydroa aestivale, the changes in the dermis are less severe than in hydroa vacciniforme. Therefore, healing takes place without scar formation.

EPIDERMOLISIS BULLOSA

In this condition resicles or bullae form usually at points of trauma but sometimes without trauma. Three forms of the disease exist epidermolysis bullosa simplex epidermolysis bullosa dystrophica and epidermolysis bullosa hereditaria letalis. The simple form is inherited dominantly, and the other two forms recessively. In the simple form the bullahe heal without scarring the mucous membranes and the nails are rarely affected and the disease improves or even sides at puberty. In the dystrophic form, the lesions heal with attrophic scars or all lesions and dystrophic changes of the nails are frequently present and the disease persists throughout life. In the third form epidermolysis bullosa hereditaria lerials death insults) occurs within the first 3 months of life. The bullae show little tendency to heal but it they heal no scars remain. Oral lesions and dystrophic changes of the nails are usually present.

Histopathology. The bullae are always in subepidermal location in the dystropluc form (Tulipan) and in the lethal form (I amb and Ealpert Schrifter) whereas in the simple form they may be found in subepidermal location (I com) in unita epidermal or in subcorneal location (Johnson and Test). However it is likely that also in epidermolysis bulloss simples the bullae always form subepidermally but because of the tendency of the epidermis to tapid regeneration the cleavage will be found intra epidermally or subcorneally in bullae that are a few days old. (This occurs not infrequently in erythema multiforme and in bullous pemphigoid see page 86). No significant inflammatory reaction is observed in the dermis unless secondary infection has occurred. In the dystroplue form small milium like epidermal cysts may be found in the upper dermis (Tulipan).

Figman and Mook first described the absence of elastic fibers in the papillary and the subpripillary layers of involved as well as non involved areas. They believed that the absence of elastic fibers was the cause of the disease Although some authors have confirmed this finding others have found in elastic cissue to be normal (Alleri). In general, it appears that in the simple form the elastic tissue is normal [Johnson and Test], while in the dystrophic form the elastic tissue in the suspension of the disease of the disease of elastic tissue in the disease of the disease tissue tends so be absent in the upper defirms of the involved areas (Tulpan Lamb and Halpert). However, it is probable that the absence of elastic tissue in the distribution by the disease process. In

46 Congenital Diseases (Genodermatoses)

HYDROA VACCINIFORME AND HYDROA AESTIVALE

Hydroa is a recurring papulovesicular eruption occurring chiefly in the summer season, usually in boys, and solely on the exposed parts of the skin. Two forms exist: hydroa aestivale, the milder form, which does not produce scarring and ends at puberty, and hydroa vacciniforme, which produces scarring and persists as a rule through-

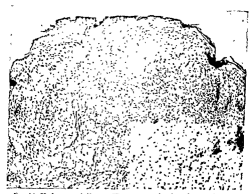


Fig. 14. Hydrox vacciniforme. There is a wedge-shaped area of necross involving the epidermis and the upper dermis. It is walled off by a chronic inflammatory inflitrate. $(\times 100)$

out life. In about one third of the cases, hydroa vacciniforme is associated with congenital porphyria, i.e., excretion of large amounts of uroporphyrin and coproporphyrin in the urine (see page 278).

Histopathology. In hydroa vacciniforme, a focal area of inflammation followed by necrosis forms in the upper detrmis (Fig. 14). In the overlying epidermis, an intra-epidermal vesicle develops which at first is multilocular, but later, because of the degeneration of epithelial cells, becomes unilocular. The vesicle is filled with fibrin, leukocytes and the débris of epithelial cells. Within the area of necrosis located in the upper dermis, the blood vessels are thrombosed and foci of hemorrhage occur. The necrotic area is walled off with a chronic inflammatory infiltrate. On absorption of the necrotic area,

ind grains (2) formation of lacunae and (3) irregular upward proiferation of papillae into the lacunae resulting in the formation of fulli (Fig. 15). There also are papillomatioss, acunthosis and hyperferations. The dermis shows a chronic inflammatory infiltrate. In



tayer are several/Morps ronds (<u>C.R.</u>), characterized by large round homogeneous deep staming nuclei. In the lower third within a <u>Machana are grant (C.</u>) characterized by small often grain shaped nuclei. (X400)

some cases in addition there is downward proliferation of epidermal rells into the dermis

The corps ronds occur mainly in the stratum malpighti and the granular liver while the grains occur mainly in the horny layer. Both may be found within the laximac Corps ronds possess large, round homogeneous deeply bisophilic nuclei and a homogeneous (hyalin ucd) deeply cosinophilic cytoplasm lined by a distinct membrane. They are much larger than normal squamous cells (Fig. 16) Corps

any case, recent studies of normal elastic tissue indicate that it plays no part in the coherence between epidermis and dermis (see page 8).

Differential Diagnosis. Differentiation of epidermolysis bullosa from other bullous diseases often is impossible. The presence of small epidermal cysts and the absence of elastic tissue, however, may aid in the establishment of the diagnosis.

DARIER'S DISCASE (KERATOSIS FOLLICULARIS)

This disease is characterized by a more or less extensive eruption consisting of hyperkeratotic or crusted papules which by confluence



Fig. 15 Darier's disease Lon magnification There arithyperkeratosis and papillomatosis Numerous <u>Leunae</u> (1) are present On the left are elongated papill limed by only one layer of cells so called yill (V) Corps ronds (CR) are present in the granular layer and grains (G) in the horag layer and in some of the facunae (X-100)

may form vertucous, crusted areas Occisionally, hypertrophic lesions are present with elevated, vertucous formations. The oral mucosa is commonly, and the vulva, the larynx and the pharynx occasionally, involved (Brunauer)

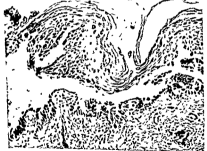
Histopathology. The characteristic changes in Darier's disease are (1) a peculiar form of dyskeratosis, namely, formation of corps ronds

tion, two rows of epithelial cells line the villi. Furthermore, no dyskeratosis occurs and large apocrine glands are present in the dermis

FAMILIAL BENIGN CHRONIC PEMPHIGUS (Hadey and Hadey)

This disease (which often, but not always, is familial) is characterized by a localized recurrent eruption of vesicles and bullae By peripheral extension, the lesions may assume a circinate conficuration.

Histopathology. Although early lesions may show like Darier's disease small suprabasal separations so-called lacunae fully devel



Fire Tr Familial brough chronic pemphigus (Hailey and Hailey). The bulls in supridusal in position One observes loss of intercellular bridges resulting in considerable acambiohism and formation of will (upward growth of papillar). These features cause a surprising resemblance to the bulls of pemphigus vilgaria (2009).

oped lessons show large separations, namely bullae in suprabasal postion (Fig. 17). With it e, elongated papillie lined by a single layer of bast cells protrude upward into the bullae and, in some cases, cords of basal cells proliferate downward into the derms. The bullae conronds develop on account of premature partial keratinization of the cell prior to reaching the horny layer, a process called benigh dys keratosis (Malignant dyskeratosis is observed in Bowen's disease and in squamous cell carcinom)

The grains are small cells, considerably smaller than the corps ronds. They resemble parakeratoric cells except that their nuclei are more prominent. The nuclei are elongated, often grain shaped

The lacunae represent small, slithke, intra epidermal vesicles which are found most commonly directly above the basal layer. They con tain desquamated, acantholytic epidermal cells which have lost their intercellular bridges due either to degenerative changes or to partial keratinization. Corps ronds and, especially, grains are present among these desquamated cells.

Elongated, often tortuous papillae lined usually with but a single layer of basal cells project into the lacunae. They often are referred to as villi

The hyperkeratosis and the pipillomatosis cause the formation of keratotic plugs. They often fill the pilosebaceous follicles but also are found outside of follicles. Darier's disease, therefore, is not, as Datter originally thought, primarily a follicular disease. Proof of this is the fact that areas devoid of follicles, such as palms, soles and oral mucosa, may be affected (Ellis). The term keratosis follicularis is, then, a mismomer.

In some cases of Darier's disease, but especially in those with hyper trophic lesions, one observes considerable downward prohiferation of the epidermis, either as a prohleration of basal cells or as pseudo epitheliomatous hyperplasia (Beerman). The prohlerations of basal cells consist of long narrow cords composed of two rows of basal cells between which there may or may not be a lacumar space. These prohlerations may send out branches and may penetrate deep mio the definits. The pseudo-epitheliomatous hyperphasic and suggest squamous cell carcinomal but, so far, no case of Darier's disease resulting in malignancy has been reported (For a discussion of pseudo-epitheliomatous hyperphasia see page 534).

The lesions on the onal and other mucous membranes are analo

The lesions on the oral and other mucous membranes are analogous to those observed on the skin except that hyperkeratotic changes are mild or absent (Brunauer)

Differential Diagnosis For the differentiation of Dariet s disease from familial benign chronic pemphigus see below. The villous pro liferations into the lacunar spaces may resemble those of syringo cystadenoma papilliferum (Beetman). However, in the latter condi-

casionally into basal cell epithelioma (Sullivan and Ellis) or squamous cell carcinoma (Costa and Junqueita; Ormea). In 1946, Lutz (one of the original describers), on the basis of suc-

In 1946, Lutz (one of the original describers), on the basis of successful implantation tests, stated that epidermodysplasia is "not an independent dermatous, but rather a generalized cruption of warts with somewhat peculiar chriacteristics" Yet, some authors, though conceding that Lutz' case was one of verrucae planae, still regard epidermodysplasia vertuciforms as an entity (Otmea: Teodorescu, Viidina) Midana obtained negative results in auto inoculation experiments performed on two patterns Teodorescu found in two patterns associated with the epidermodysplasia vertuciforms, kera toosy palmants et plantaris hereditatia and regarded both diseases as genodermatoses.

ACROKFRATOSIS VFRRUCIFORMIS (Hopf)

Numerous hyperkeratotic and occasionally vertucous papules are present, predominingly on the doisy of the hands and the feet

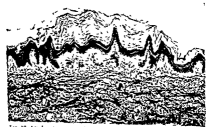


Fig. 18 Acroleratosis terruciformis (Hopt). The lesion consists of a circum scribed area of papillomatous, acanthosis and hoperheratosis. (x100)

Histopathology. The papules show considerable hyperkeratosis, increase in thickness of the granular layer and acanthosis (Fig. 18) In addition, there are slight papillomatosis and some thickening of the rete ridges. The rete ridges all extend to a uniform level and their configuration is well preserved. There is no parakeratosis or

E3

tuin acontholytic cells some of which show evidence of keratinization Occasionally there is evidence of dyskeratosis namely shrinking of some of the acontholytic cells which then assume the appearance of grains. However corps ronds have been described in only a few cases (Ellis Winer and Leeb)

Differential Diagnosis Familial benign chronic pemphigus shares certain histologic features with both Darier's disease and pemphigus vulgaris. In all three diseases one finds suprabasal separations of the epidermis caused by acantholysis and upward proliferation of papillae as so called villi into the resulting lacture or bullae. I amilial benign chronic pemphigus however differs from Darier's disease by the larger size of the suprabasal separations (which thus appear as bullae rather than as lacture) and the lesser degree or even the absence of dyskeratosis. If dyskeratosis is absent differentiation from pemphigus vulgaris may be impossible Occasionally however one observes in familial benign chronic pemphigus even in the absence of dyskeratosis. In early onset of keratinization in the lower layers of the detached epidermis. The presence of cosinophils in the bullae points toward a diagnosis of pemphigus vulgaris but their absence does not rule it out.

The nosologic position of the disease is at present uncertain. The presence of dyskeratotic changes in most cases of familial being chronic pemplinguis has led some observers to regard this disease as a bullous variou of Datier's disease (Ellis Finnerud and Szymanski Winer and Leeb). However, until more is known about the cause of the disease, it may be best to regard it as an independent entity

(Huley and Huley Frank and Rein)

EPIDFRMODYSPLASIA VERRUCIFORMIS

(Lewandowsky and Lutz)

In this derimatosis one finds an extensive eruption of flat topped hyperkeratotic lesions resembling verrucae planae B₃ confluence

lichenified plaques may form

Histopathology The histologic changes consist of hyperkeritosis increase in the thickness of the granular layer acanthosis and a peculiar vacuolization of the cells in the upper layers of the stratum malpighin of the granular cells and of the horns cells Because of the vacuolization the horns layer shows a loosely felted basket weave pattern. The histologic picture is like that of vertuca plana (Fig. 122) with two exceptions (1) in epidermodysplasia vertucal formis the nuclei of the vacuolited cells show more pronounced pyknosis and frigmentation than in vertuca plana (Sullivan and Ellis Waisman and Montgomery) and (2) the lesions change of

casionally into basal cell epithelioma (Sullivan and Ellis) or squit

mous cell carcinoma (Costa and Junqueira, Ormea) In 1946, Luiz (one of the original describers), on the basis of successful implantation tests, stated that epidermodysplasia is "not an independent dermatosis, but rather a generalized cruption of warts with somewhat peculiar characteristics." Yet, some authors, though conceding that Lutz' case was one of vertucar planae, still regard epidermodysplasia vertuciformis as an emity (Ormea, Teodorescu, Midana) Midana obtained negative results in auto inoculation experturents performed on two patients Teodorescu found in two patients, associated with the epidermodysplasia vertuciformis, keratosis palmaris et plantaris hereditaria and regarded both diseases as genodermatoses

ACROKERATOSIS VI RRUCIFORMIS

(Hopf)

Numerous hyperkeratotic and occasionally verrucous papules are present, predominantly on the dorsa of the hands and the feet



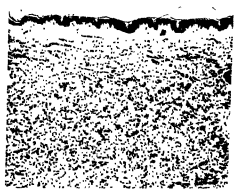
Fig. 18 Actokeratosis vertuciformis (Hopf). The lesion consists of a circum scribed area of papullometers, accombine and hyperkeratosis (X100)

Histopathology. The papules show considerable hyperkeratosis, increase in thickness of the granular layer and acanthosis (Fig. 18) In addition, there are slight papillomatosis and some thickening of the rete ridges. The rete ridges all extend to a uniform level and their configuration is well preserved. There is no parakeratosis or

vacuolization of cells such as is seen in verruca vulgaris and verruca plana (Loveman; Niedelman).

PSEUDOXANTHOMA ELASTICUM

This disorder represents a congenital defect of the elastic tissue which may be limited in extent or widespread. In addition to the elastic tissue in the dermis, the elastic membrane of the retina, called



Pseudoxanthoma elasticum, Low magnification, Verhoeff stain. This illustration shows the usually observed late degenerative stage. In the lower dermis, the clastic fibers are increased in number, appear swollen and show signs of degeneration, such as fragmentation, disintegration and clumping (Patient's age, 31 years) (×50)

Bruch's membrane, and the elastic fibers in the aorta, the arteries and the arterioles may be affected. The degeneration of the elastic fibers of Bruch's membrane causes small defects and breaks which manifest themselves clinically as angioid streaks (Urbach and Wolfram, Hagedoorn, Ebert) Involvement of the aorta causes diffuse dilatation (Urbach).

The cutaneous lesions consist of soft, yellowish papules and plaques The papules frequently show a linear arrangement. The sides of the neck are the most common site of the lesions

Histopathology. Histologic examination of the skin reveals the elastic tissue to be normal in the subepidermal portion of the dermis

In the middle and lower portions of the dermis, however, it is in creased, usually in circumscribed areas. Within these areas, the elanic fibers are swollen and show

tion, disintegration and clu

eration and imbibition with a basophilic staining and become visible with routibe stains. Fin

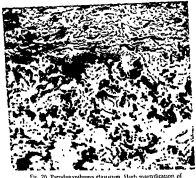


Fig. 20 Pseudoxanthoma elasticum. High magnification of Liguise 19 terhoeff stain. The elastic fibers are degenerated whereas the collagen shows no evidence of degeneration $(\times 200)$

nerud and Nomland have shown by staming methods (von Kossa's stain for calcium) and Lobitz and Osterberg by micro incineration that the degenerated elastic fibers are righly infiltrated with calcium The collagen fibers are unaltered

In a few instances, the elastic tissue of the arterioles and the small arreries in the deep demus was found to have undergone a similar degeneration (Urbach, de Si Penella and Esteves)

It appears that pseudoxanthoma elasticum occasionally is preceded by an early hyperplastic stage (Fig. 21). In this stage, which seems to be present only during infancy and childhood, the elastic fibers are increased in number and size but show no evidence of degeneration as they do in the more commonly observed late degenerative stage (Weidman, Anderson and Ayres, Pautrier)

Differential Diagnosis. Senile elastosis, like pseudovanthoma elasti cum, shows a great increase in material taking the elastic tissue stain However, in senile elastosis this material is located in the upper third



Fig 21 Pseudovanthoma elasticum Los magnification, Ver hoeff stain. This illustration shows the early hyperplastic stage. The elastic fibers are large and shollen but show no evidence of degeneration (Patient's age, 2 years) (×100).

of the dermis and is present as dense masses rather than as individual curls. Furthermore, staining of this material for calcium is always negative in semile elastosis.

CUTIS HYPERELASTICA (EHLERS DANLOS SYNDROME)

This syndrome consists of (1) hyperelasticity of the skin, (2) hyper extensibility of the joints, (3) fragility of the skin and the blood vessels with formation of atrophic scars and (1) development of raisin like pseudotumors. The pseudotumors form at points of trauma and are soft and pigmented and present a wrinkled surface.

In many cases small, hard, subcutaneous nodules have been described They also are the result of trauma.

The collagen bundles appear atrophic, split up and separated by edema. The clastic fibers are normal in some cases but in most cases show breaking up and clumping. The amount of elastic tissue often appears increased but it is likely that this increase is not a real one but is simulated by the atrophy of the collagen. In addition, the number of capillaries is increased and their lumina dilated. Large, round cystic spaces representing lymphangiectatic civities may occur (Korting and Gottron)

The raisinlike pseudotumors that are part of the syndrome form at areas of traumatic hemorrhage and consist either of accumulations of foreign body giant cells (Ronchese) or of proliferated connective nodules contain calcified necrotic fat or mucoid material enclosed in a thick fibrous capsule (Johnson and Falls)

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URTICARIA PIGMENTOSA

This disorder is characterized in most cases by the presence of a great number of brown macules scattered over the entire curaneous surface When tubbed with a blunt instrument, they become dis uncily urticarial In rare instances the lesions consist of one or several soft nodules or plaques

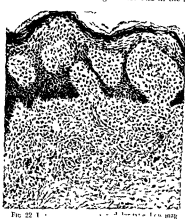
Historiathology The histologic picture shows an infiltrate composed chiefly of mast cells. Mast cells are characterized by the presence of basophilic metachromatic granules in their cytoplasm (see pages 37 38; These granules are not visible on staining with routine st uns. Their demonstration requires special staining [see pages 28] 29 and Plate I) Fixation in 10 per cent Formalin (without subse quent exposure to Helly's solution) and staining with Giemsa's stain or with methylene blue are recommended

In the micular type mast cells are present in the upper third of the dermis especially around the capillaries Some appear to be round or oval but the majorny are spindle-shaped. Since in sections stained with hematoxylin and eosin the mast cells resemble fibro blasts the diagnosis may be missed easily unless special staining is employed

In the nodular type (Figs. 22-23) mast cells lie closely packed in tumorlike aggregates. The infiltrate may extend through the entire dermis into the subcutis. Wherever the mast cells he in dense aggre gates they are cuboidal rather than spindle shaped and show ample, slightly cosmophilic cytoplism Because of their shape and ample 58

cytoplasm, they look unlike any other cell and the drignosis can be made without resorting to special staining

If a biopsy is performed shortly after the lesion has been stroked, the section will show edema, an influx of eosinophils and shrinking of the mast cells associated with a great decrease in the amount of

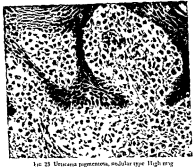


nification packed in Because of staining with hem novelin cosin, the granules in the mist cells are not visible (×200)

granules within them indicating that the granules have been expelled from the cells (Drennan) There may even be disintegration and a temporary disappearance of may cells which may be the explanation for certain reports in the literature where in otherwise typical cases of urticaria pigmentosa mast cells have not been found (Drennan and Beare)

The pigmentation of urticaria pigmentosa is due not to the pres ence of the most cells but to melanin which is present in increased amounts in the basal layer and occasionally also in melanophores

The presence of extracutaneous lesions in utilicaria pigmentosa has been reported recently However, the systemic lesions in Ellis' case may well have been lymphoma, and the bone lesions in Sagher's and Clyman's cases were not examined histologically



milication of Figure 22 hematoxylin cosin stain The mast cells appear as large cuboidal cells (X400)

INCONTINENTIA PIGMENTI

The disease frequently begins immediately after birth with inflammatory lesions, particularly bullae, in linear and grouped arrangement The bullae may recur for months They finally give way to areas of pigmentation. There may be an intermediary stage of linear vertucous lesions. The pigmented areas are widely dissemi nated, located especially on the trunk, and have an irregular, bizarre outline

Histopathology. In the early vesicular stage, one observes intraepidermally located vesicles containing many eosinophils (Carney, Fpstein, Vedder and Pinkus) In addition, there is spongiosis of the epidermis, and in the dermis an infiltrate composed of lymphon to eosinanh le an 1 -

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The final stage shows diminution or absence of melanin in the basal laver of the epidermis and extensive deposits of melanin inside

eytoplasm, they look unlike any other cell and the diagnosis can be made without resorting to special staining

If a biopsy is performed shortly after the lesion has been stroked, the section will show edema, an influx of eosinophils and shrinking of the mast cells associated with a great decrease in the amount of

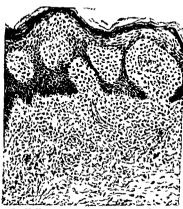


Fig. 22 Urticaria pigmentosa, nodular type Low mag infication hemitoxylin eosin stain. Mast cells he closely packed in the upper dermis. They are cuboid if in shape Because of staining with homotoxylin cosin, the granules in the mist cells are not visible (×200)

granules within them indicating that the granules have been expelled from the cells (Drennan) There may even be disintegration and a temporary disappearance of most cells which may be the ex planation for certain reports in the literature where in otherwise typical cases of urticaria pigmentosa mast cells have not been found (Drennan and Beare)

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and outside of melanophores in the upper dermis. In many cases the basal cells show degeneration and vacuolization. It is believed generally that the disease causes damage to the cells in the basal layer so that the melanocytes become incapable of holding and metabolizing melanin (Sulzberger, Doornink).

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TABLE 2 -CLASSIFICATION OF VESICLES AND BULLAE

		n +=====	
TYPE OF VESICLE OR BULLA	MODE OF FORMATION	SITE OF FORMATION	Diseases
Subcorneal bulla	Detachment of horny Javes	Subcorneal	Impetigo
Spongtotte bulla	Cellular necrosis fol lowed by spongiosis and, occasionally, by reticular degen eration	Intra epidermal	Dermatius-Ecrema Pompholys Congenital ichthyosiform erythroderma Hydroa
Mihamal bulla	Retention of sweat	Intracorneal Intra-epidermal or subepidermal	Miliana crystallina Miliana rubra
Acantholy tie bully	Acantholysis	Intra epidermal a suprabasal	Pemphigus vulgans Pemphigus vegetans Familia lenging chronic pemphigus Daner's disease Senile keratosis
		b upper epi	Pemphigus foliaceus Pemphigus erythematosus
5 Viral bulla	Reticular and ballooi ing degeneration with acantholysis		Vanola Herpes simples Herpes zoster Vancella
6 Pressure bulla	Detachment of entir epidermis		Bullous pemphigoid Benign mucous mem- brane pemphigoid Dermatitis herpetiformis Erythema multiforme Epidermolysis bullosa Porphyna cutanea tarda
 Bulla due to base cell degeneration 	crills		Incontinentia pigmenti Lichen planus Lichen sclerosus et atrophicus Lupus erythematosus
8 Balla due to reta lum degeneratio		b Subepidermal	Burns
			

7

Noninfectious Vesicular and Bullous Diseases

Several new concepts have been introduced in recent years concerning the histologic structure of vesicles and bullae and their mechanism of formation. It appears appropriate therefore to present a classification of the different types of vesicles and bullae and to outline briefly their mode of formation before discussing individual diseases (Since from a histologic point of view it is immaternal whether a lesion is a vesicle or a bulla-only the latter term will be used in the following presentation.)

Light distinct types of bullae can be recognized (Table 2)

I Subcorneal bulla Detachment of the horny layer from the stratum malpighu occurs

2 Spongiotic bulla Liquefaction necrosis of a few epidermal cells causes formation of a minute causy (icisculette primordiale) which subsequently due to intercellular edema (spongiosis) and intracellular edema (alteration cavitaire) in the surrounding epidermis en larges into an intra epidermal bulla. In the case of pronounced intra cellular edema reticular degeneration of the epidermis may occur (For a detailed description see page 69).

3 Miliarial bulla In miliaria bullae form due to the escape of sweat from the sweat ducts. Two types occur miliaria vrystallina in which the bulla is locited within the stratum corneum and miliaria dubra, in which the bulla forms either within the stratum malpighi or beneath the epidermis. In severe cases of miliaria rubra, the sweat duct ruptures. (For a detailed description see page 76)

4 Acantholytic bulla Degeneration of intercellular bridges causes loss of coherence between epidermal cells and formation of rifts which enlarge into bullae Detached (acantholytic) epidermal cells are present in the bulla cavity. Acantholysis may take place within the lower epidermis predominantly right above the basal layer or within the upper epidermis predominantly within the granular layer (For a detailed description see pages).

TABLE 2 -CLASSIFICATION OF VESICLES AND BULLAE

		A	
Type of Vesicle	Mode of Formation	91TE OF FORMATION	Diseases
Subcorneal bulla	Detachment of horny layer	Subcorneal	Impetiga
Spongiotic bulla	Cellular netrosis fol lowed by spongrous and, occasionally, by reticular degen eration	Intra-epidermal	Dermatitis-Eczema Pompholyx Congenital schthyosiform erythroderma Hydroa
Miliarral bulla	Retention of sweat	Intracorneal Intra epidermal or suhepidermal	Mihana crystallina Mihana rubra
Acantholytic bulls	Acantholysis	Intra-epidermal a suprabasal	Pemphigus vulgaris Pemphigus vegetans Familial benign chronic pemphigus Daner's disease Senile keratosis
		6 upper epi dermis	Pemphigus foliaceus Pemphigus erythematosus
Viral bulia	Reticular and balloon ing degeneration with acantholysis	Intra epidermal	Vanola Herpes simplex Herpes zoster Vancella
S Pressure buils	Detachment of entire	Subepodermal	Bullous pemphigoid Benign mucous-mem brane pemphigoid Dermatitis herpetiforms Erythema multaforme Epidermolysis bullosa Porphyria cutanea tarda
7 Bulla due to basa cell degeneration	Degeneration of base cells	Subepidermal	Incontinentia pigmenti Lichen planus Lichen sclerosus et atrophicus Lupus erythematosus
8 Bulla due to retic		b- Subepidermal	Burns

5 I tral bulla Invasion of epidermal cells by certain viruses causes 5 I trat butta Invision of epidermal cells by certain viruses causes two types of degenerative changes in epidermal cells ballooning and reticular degeneration. Ballooning degeneration leads to extensive acantholysis. Although the bullar form within the epidermis progression of the ballooning degeneration leads to a subepidermal location in older bullar. (For a detailed description see page 239).

6 Pressure bulla. Detachment of the entire epidermis from the dermis occurs. Pressure of the accumulating tissue fluid often causes.

in early bullie rounding of the literal wills and stretching of the epidermal cells located there Older bullie due to regeneration of the epidermis at the floor of the bulla may have an intra epidermal location If in addition disintegration of the detached stratum malpighii occurs the bulla may be subcorneal in location (For a

detailed description see page 86)
7 Bulla due to basal cell degeneration Several diseases (see Table 2) cause hydropic degeneration of the basal cells. This may result in damage to the cytoplasmic processes which effect the coherence of the basal cell layer with the dermis. Thus, a subepidermal bulla may

form (For a detailed description see page 296)

8 Bulla due to reticulum degeneration Damage to the subept dermil feltwork of reticulum fibers causes the cytoplasmic processes of basal cells to pull out of this feltwork. A subepidermal bulla forms into which numerous cytoplasmic processes extend from the basal cells at the roof of the bulla (For a detailed description see page 00.1

DERMATITIS ECZEMA

The terms dermatitis and eczema now are used generally as syno nyms They refer to n inflammation of the skin based on an allergic response of the skin to a variety of agents such as chemicals proteins bacteria and fungi. The exciting illergen may act on the skin either from the outside or from the inside

Dermatitis or eczema may be acute subacute or chronic The clinical picture is characterized by polymorphism of the eruption Among the primary lesions that may be observed are macules Among the primary lesions that may be observed are micules papilles and vesicles the macules and papilles tend to coalesce to form areas of diffuse crythema. Among the secondary lesions are scaling crusting lichenification and fissuring. The lesions of dermatitis usually are not demarcated sharply but merge gradually into the surrounding normal skin. Moderately severe to severe atching is present in most forms of dermatitis. No generally accepted hassification of dermatitis exists and many cases dely assignment to any definite type. In this section, the following types of dermatitis will be discussed. (1) contact dermatitis

(2) nummular eczema (3) atopic dermatitis (or neurodermatitis dis seminata) (4) lichen simplex chronicus (or neurodermatitis circum scripta) (5) extidative discoid and lichenoid chronic dermatosis (6) scripts) (3) extuative checking the interior of notice that sees (7) starts derinativis and (8) generalized exfoliative derinativis in addition fessions of derinativis may occur in superficial fungus infections as drug eruptions and in lymphoma. The latter three forms of dermatitis will be discussed when the respective diseases are described

Contact derinalitis is caused by contact of the skin with an agent

that acts either as a specific allergic sensitizer or as a primary irritant Contact dermatitis may be acute, subacute or chronic. In acute and subacute contact dermatitis diffuse erythema edema oozing and crusting predominate in addition vesicles are often present if a specific allergic sensitizer is the cause. In chronic contact derinatitis erythema scaling and lichenification prevail

'ummular ec ema probably caused by a temporary loss of re sistance in the skin to the ordinary bacterial flori of the skin presents fairly sharply demarcated patches of erythema studded with discrete

pinpoint vesicles or pinpoint erosions

Atopic dermatitis a constitutional familial dermatosis of unknown cause which often is aggravated by emotional tension or allergic factors shows inhemified and scaling erythematous areas which when netive show also oozing and crusting but no vesicles

Lichen simplex chronicus shows one or several lichenified plaques with little scaling. Oozing and vesiculation are absent. On the lower legs especially lichen simplex chronicus may become hypertrophic and assume a verrucous nodular appearance (lichenificatio gigantea lichen corneus hypertrophicus)

Exudative discoid and lichenoid chronic dermatosis short s me h lesions wf

and later sent an er

- - a or nummular eczerna

Seborrheic dermatitis shows fairly sharply demarcated brownish red areas which show only little infiliration and are covered with ted areas which show only more infinition and are covered with hine greasy scales. Oozing may be present but no vesiculation is found Generalized seborrheic dermatitis in infants often is referred to as Leiner's disease

Stasis dermatitis presents erythemy edema scaling and occasionally cozing and crusting. It differs from other forms of dermatitis first by showing brownish pigmentation due to hemosiderin denos

oozing. It represents a peak reaction to which several forms of derma titis may lead—for instance, contact dermatitis, atopic dermatitis, seborrheic dermatitis, stasis dermatitis, drug dermatitis and lym phoma dermatitis. However, it may occur as an idiopathic disease

Histopathology. The various types of dermatitis rarely present a histologic picture sufficiently diagnostic to allow their differentiation



Fig. 21 Acute dermaints contact dermains due to poson sy. Nu merous intro-epidermilly located vesicles and myrked introcellular democrate present. The vesicles are separated by thin septi formed by the resisting wills of edemicors epidermil cells (reticular degeneration) and thus form a multilocalir bulls. (2016)

because the same histologic reactions occur in all forms of dermatitis exidation leading to vesiculation in the acute stage, proliferation leading to acanthosis in the chronic stage, and a combination of these two reactions in the subrcute stage. Since, as a rule, no more specific diagnosis than acute, subacute or chronic dermatitis can be made, the histologic picture as presented by an acute, subacute and chronic dermatitis will be described first. Thereafter, the distinctive features occasionally presented by the various members of the dermatitis eczema group will be discussed.

IN ACUTE DERMATITIS, intra epidermally located esicles or bullae predominate the histologic picture. Considerable intercellular edema (spongiosis) and intracellular edema (altération cavitaire) may be present in the epidermis surrounding the vesicles. If the number of

vesicles is great and the intrucellular edema pronounced the vesicles due to reticular degeneration of the epidermis will be separated from one another only by thin septa formed by the resisting walls of edematous epidermal cells and will thus form a multilocular bulla (Fig 24) The vesicles and the bullae contain a few lymphocytes cosmophils and neutrophils and disintegrated epidermal cells vilgrating lymphocytes and neutrophils are present in the epidermis. The cells of the stratum corneum may be parakeratoric and intermingled with fibrin and numerous neutrophils (Viner and Liptungled with fibrin and consistence).

neuroj // Formation of the vesicles and the bu

11 had been 1e vesteles necrosis is

the primary factor and that spongroup / In 1925 Creatte described as the primary lesion the vesiculette primordiale formed by the lysis of two or three squamous cells through cyto

formed by the lysis of two or three squamous cells through cyto plannic alteration. Spongiosis followed this and caused enlargement to a expressed the same

ipport vesicles tend

to form in areas where the rete tets appear of effect vesicles often he in areas formerly occupied by epidermal cells (Fig. 25) and spongious may be entirely absent in the vicinity of vesicles Polak and Mom studying the formation of vesicles in experimental ecrema with the aid of silver impregnation found that at the time the vesiculette primordiale formed the intercellular bridges were still intact and only broke later by mechanical force when the vesicle had integrated in size

Differentiation of acute vesicular or bullous dermaitis from bul louserythema multiforme and dermatitis herpetiformis is not always possible Although in the latter two diseases the bullae form subepi dermally they may be located intra epidermilly during the stage of healing due to regeneration of the epidermis. In that case secondary findings such as the presence of marked spongiosis in contact dermains and its absence in the other two diseases and the number of cosmophils may aid in the decision. It should be stressed as of utmost importance that for the diagnosis of all vesicular and bullous diseases an early lesion must be chosen for histologic examination be cause eccondary factors such as regeneration and pyogenic infection may obscure the diagnosis of the properties of the properties of the diagnosis of the properties of t

outing. It represents a peak reaction to which several forms of derma titis may lead-for instance contact dermatitis atopic definatitis seborrheic dermititis stisis dermititis drug dermititis and him phoma dermatitis. However it may occur as an idiopathic disease

Histopathology The various types of derinatitis rarely present a histologic picture sufficiently diagnostic to allow their differentiation

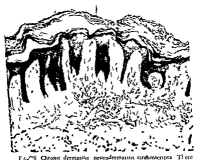


Fig. 24 Acute dermatitis contact dermatitis due to poison in Nu merous intracepidermally located vesicles and marked intracellular edeur, are present. The vesicles are suparited by thin septa formed by the resisting wills of edentitious epidermal cells (reticular degeneration) and thus form a multilocular bulla (×100)

because the same histologic reactions occur in all forms of dermatitis exudation leading to vesiculation in the acute stage proliferation leading to wanthosis in the chronic stage and a combination of these two reactions in the subscure stage. Since, as a rule no more specific diagnosis than acute subacute or chronic dermatitis can be made the histologic picture as presented by an acute subscute and chronic dermittus will be described first. Thereafter the distinctive features occasionally presented by the various members of the derma titis eczema group will be discussed

In acute permatitis intra epidermally located esicles or bullae predominate the histologic picture. Considerable intercellular edema (spongiosis) and intracellular edema (alteration cavitaire) may be present in the epidermis surrounding the resicles. If the number of

Sachs Muller and Gray (1946) to the histologic picture as described above for chronic derinatitis because it is characteristically present in neurodermatitis disseminata (atopic dermatitis) and neurodermatitis circumscripta (lichen simplex chronicus). However the same histologic picture may occur in any dermatiosis belonging to the dermatitis eczema group.



F 6,7% Chronic dermatutes neurodermatus cargametripts. There are their transitions acanihos a clongar on of the receivings and clongar to no and broadening of the pap line. The often a sho s a chron c in flummatory infiltrate and fibros s (X50).

Many diseases not members of the dermatics eccent group show either regularly or occasionally a histologic picture allowing no more specific diagnosis than chronic dermatitis Diseases which regularly show, the nonspecific histologic picture of chronic dermatitis methodomy, cause roses parapsoriasis and pellagra. Many other discusses such as psoriasis, lichem planus and luquis erythematosis to native but a fen show a diagnosiic histologic picture in clinically ity is al cless but may show a nonspecific histologic picture that of chronic dermatitis in clinically at pical cases.

I relycosis fungoides must always be kept in mind as a possible diagn ses when a section showing chronic dermatitis is examined. It often is a very difficult task to establish or rule out early mycosis In subacutt permatitis, one sees pongiosis, intracellular edema and, usually also esicle formation. However, the vesicles are smaller than in acute dermatutis (Fig. 25). Moderate a canthosis and varying degrees of parakeratosis are present. The inflammatory infiltrate in the dermis usually is pronounced and is composed of a multiplicity of cells. Symphocytes predominate, but neutrophils, gosinophils and



Fig. 25 Subacute dermatitis nummular eczema There is intra epidermal (essele lormation The vesteles lie in areas formerly occu pied by epidermal cells /The epidermis shows, parkertiesis and mod erate acauthosis. The vermis shows a peris vegular infilitate. (X100)

 Instrocytes are also seen. There may be considerable migration of neutrophils and lymphocytes through the epidermis

In Chronic derivativis, there is often marked acanthosis with elongation of the rete ridges. There is hyperkeratosis intermingled with areas of parakeratosis. Slight intercellular edema may be present in the epidermis, but vesicle formation is absent. In the upper dermis, one sees a moderate amount of predominantly pervastular infiltration composed of various types of cells. Lymphocytes prevail, but the number of eosinophils, histocytes and fibroblasts may be considerable. Neutrophils are absent. The number of capillaries is micreased and the walls of the arterioles and the small arteries may be thickened. The term neurodermatitic reaction has been given by

that of psorrsis which shares with lichen simplex chronicus the tendency to elongation of the rete ridges. However, psoriasis shows predominantly partieratosis rather than hyperheratosis thinning of the suprapapillar, portions of the stratum milyighti edemy of the upper portions of the papillae and not infrequently. Murror micro abscesses Furthermore the papillary capillaries in psoriasis are diated and tortuous whereas in lichen simplex chronicus they appear normal According to Stoughton and Wells the Hotchkiss McManus and descriptions. normal Accounting to brighters than seem psoriasis very clearly and thus aids in its differentiation from lichen simplex chronicus

thus aids in its differentiation from fichen simplex chronicus FXUDATIVE DISCOID AND LICHENOID CHRONGE DERMATOSIS (Sulz berger Garbe) The epidermis may show spongiosis and vesiculation as in subacute dermatitis or acanthosis with little or no edema as in chronic dermatitis. The vessels of the upper dermis and the mid dermis are dilated and their walls thickened. About them is a mantle of lymphocytes histocytes polymorphonuclear neutrophils cosino-phils and numerous plasma cells Sachs and Kirsch state that the pres ence of many plasma cells gives the histologic picture a distinctive appearance so that the diagnosis can be established on microscopic findings alone

SEBORRHEIC DERMATITIS The histologic picture is not diagnostic SERORITEE DEVANTITY The histologic picture is not diagnostic it may be said to be halfway between psortiasis and chronic dermantis. The horn Jayet because of the tendency to desquamation is only poorly developed and most of its cells are parakeratotic. The epi detrims shows slight to moderate acambous with elongation of the reterridges and slight intracellular edema and spongouss. The detrims shows a mild chronic inflummatory infiltrate. Munro micro abscesses and neutrophils migrating through the epidermis as seen in psoriasis occasionally are observed. In cases in which the histologic picture

resembles that of psorruss the presence of spongos s car a

on unimis Older lesions may show numerous newly formed capillaries embedded in a fibrotic dermis Whereas it has been assumed generally that venous and capillary stasis is responsible for the chinical manifestations of stasis derma titis kulum and Hines in a study of the vessels at the dermo-sub cutaneous junction found changes more often and more severely in the arterioles than in the venules. The changes in both arterioles and venules consisted of intimal proliferation endothelial hyper-plasia and medial hypertrophy Complete obliteration of atterioles

Noninfectious Vesicular and Bullous Dispases

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fungoides One should search for attpical instincties (so-called my cosis cells) mitotic figures clumping of nuclei haryorthesis (disinte gration of nuclei into nuclear dust) and <u>Paintier micro</u> obscesses (For further details see page 485) However it should be realized that some atypicality of the histocytes and an occasional mitotic figure sometimes may be seen in chronic detrinitis <u>Handoubt, it is always</u> best to request another specimen for histologic examination

A few words about the histologic aspects of the various members

of the dermatitis eczenia group are now in order

CONTACT DERNATITIS Contact dermatitis may be acute subacute or chronic. The histologic descriptions given above for acute subacute and chronic dermatitis priging in general to contact dermatitis. Acute contact dermatitis presents numerous closely set large and small intra epidermal vesicles (Fig. 24). Chronic contact dermatitis shows irregular acanthosis. Even at that stage spongiosis and small intra epidermal vesicles are often present (Sachs Miller and Gray 1914. Miller).

NUMBULAR FCZENTA (infectious eczematord dermatitis). This eruption characterized clinically by pinpoint vesicles usually shows histologically a picture of subactite dermatitis (Fig. 25). In a moderately acunthotic epidermis one finds numerous scattered areas of intra epidermal vesiculation (Sachs, Miller and Gray, 1946). As a rule spongiosis about the vesicles is slight or absent.

Atolic Dermatitis (neurodeimatitis disseminata) The histologic picture is usually that of a chronic dermatitis showing acanthosis with varying degrees of spongrosis. The number of eosinophils in the inflammatory infiltrate is often considerable (Burkhart and Montgomery).

Lichen Simites Chronicus (neurodermitius cucumscripti) The microscopic appearance is essentially that of a chronic deminitis (Fig 26) There are hyperkeratosis interspersed with small areas of parakeratosis acanthosis characterized by rather regular elongation of the rete ridges and elongation and broadening of the papillae. There may be some spongiosis but vesiculation does not occur. In addition to a chronic inflammatory infiltrate the dermis often shows a fur number of fibroblasts and some fibrosis even in the papillae.

In the hypertrophic type of lichen simples chronicus (lichentificatio gigantea or lichen corneus hypertrophicus) the epidermis shows in addition to acanthosis with elongation of the rete ridges considerable hyperkeratosis and papillomatosis (Shaffer and Beerman Hymra and Erger)

The histologic picture of lichen simples chronicus may resemble

tion at first shows no evidence of lymphomatous infiltration in the dermis it is advisable to perform further biopsies at intervals (See also page 490)

DERMATOPATHIC LYMPHADENITIS

Any extensive dermaturs but particularly generalized exfoliative dermaturs whether due to lymphoma or not may cause a generalized lymphademus of the subcutaneous lymph nodes. Histopathology This lymphademus has certain characteristic his tologic features not found in other types. It was described first by Pautrier and Woringer as hipomelmour teticulosis. Hurwitt introduced to the subcutage of t

hmph rminal

centers. The pulp of the lymph node shows considerable hyperplasia offreticulum cells. Since the reticulum cells possess abundant and offreticulum cells Since the reticulum cells possess abundant and caintly cosinophilic cytoplasm the areas of reticular hyperplasm appear as large pale patches (Laipply). The reticulum cells show phagocytic activity and may control hemosiderin melanin and, occasionally, fair. The lymph follicles as well as the pulp of the lymph node are permeated with cosinophils neutrophils and plasma cells. The intermediary sinuses are filled with reticulum cells. (sinus catarrh) The melanin and the fat occasionally present in the lymph nodes originate in the skin and are carried into the lymphatics by scratching It has been suggested that the fat may not be sebum but ointment base (Bettley)

Differential Diagnosis These histologic changes differ from those observed in mycosis fungoides. Hodgkin's disease and follocular lym phoma by the absence of destruction of the basic architecture of the lymph node the absence of Sternberg Reed cells and the pres ence of phagocytic activity in the reticulum cells (Hurwitt) The large lymph follicles present in dermitopathic lymphadenitis differ from those of follicular lymphoma by greater uniformity in size smaller number and absence of fissures which frequently separate the lymph follicles from the stroma in follicular lymphoma

In recent years the relationship of dermatopathic lymphadenitis to ismphoma has been discussed by several authors. Some have assumed that a dermatopathic lymphadenitis can develop into lym phoma (Bluefarb and Webster) and others have regarded cases of ceneralized erythroderma with dermatopathic lymphadenitis as Brill Symmers disease (i.e. follicular lymphoma) even in the absence of the histologic criteria of follocular lymphoma (Rost) Neither point of view is justified Dermatopathic lymphadenitis as such is an en

was seen in specimens from ulcerated areas. These authors concluded

that arteriolai changes may be an important etiologic factor in stasis dermatitis GENERALIZED EXPOLIATIVE DERMATITIS The histologic appearance may be that of a subacute or a chronic dermatitis. In the subacute

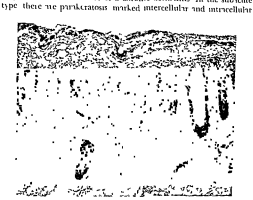


Fig 27 Generalized exfoliative dermatitis due to arsphenamine There are parakeratosis marked intercellular and intracellular edema in the upper stratum malpighii acanthosis with elongation of the rete ridges and migration of inflammatory cells through the epidermis. The upper dermis shows edema and a considerable amount of inflammators infiltrate (×100)

edema, particularly in the upper stratum malpighic acouthosis with elongation of the rete ridges and migration of cells through the epi dermis The upper dermis shows edema and a considerable amount of inflammatory infiltrate (Fig. 27). If the edema in the upper stratum malpighii is pronounced the cells of the upper stritum may exfoli ate together with the parakeratotic horns cells. In the chronic type of exfoliative dermatitis the histologic picture is that of chronic der matitis. Each such case requires thorough histologic investigation in order to rule out lymphoma Montgomery has stated that 25 per cent of all cases of exfoliative dermatitis are proved on histologic examina tion to be associated with lymphoin? Even if the histologic examina

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Erythema Multiforme

Costello M J Erythema multiforme exudatisum J Invest Dermat 8 127 1947 Winer L. H. and I ipschultz C. E. Comparative study of histology and extology in vesiculating eruptions Arch Dermat & Syph 65 270 1952 Burn

Moritz & R. Studies of thermal injury. III. The pathology and pathogenesis of cutaneous burns Am | Path 23 910 1947

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tirely nonspecific reaction to an extensive dermatitis. Since lymphoma may manifest itself as an extensive dermatitis (see page 490), it can cause a dermatopathic lymphadenitis. Subsequently, the lymphoma may extend to lymph nodes previously affected by dermatopathic lymphadenitis, but in that case the latter is only chronologically—not etiologically—the foretunner of the lymphoma (Jarrett and Kellett; Keller and Staemmler)

MILIARIA

Miliaria occurs following excessive sweating in parts of the body covered by clothing. There are two types miliaria crystallina and miliaria riibra. In miliaria erystallina, asymptomatic, small, super ficial, noninflummatory vesicles are present. In miliaria riibra, the lesions consist of pruritic, discrete but closely aggregated papules papulovesicles and resicles surrounded by erythema.

The cause of miharia lies in excessive hydration of the horny layer by sweat. This results in swelling of the keratin, closure of the narrow sweat pores by keratin and retempon of sweat in the sweat ducts.

Histopathology. In miharia rystallina, histologic examination reveals occlusion of the orifices of sweat ducts by keratin plugs and distention of the sweat ducts within the epidermis and, occasionally also in the dermis Intracorneal vesicles are present. There is notin flammatory infiltrate. Evidence in favor of the assumption that the intracorneal vesicles contain sweat are the observations in experimental miharia crystallina by Shelley and Horvath that the vesicles on serial sections proved to be in direct communication with sweat ducts and failed to form when sweating was inhibited by the local intection of attripine.

In miliaria vibra an inflammatori, infiltrate is present around the sweat ducts in the epidermis and the upper dermis. Either an intra-epidermal or a subepidermal vesicle is seen. It appears that in severe cases of miliaria rubra, which are associated with thermogenic anhi-drosis, the sweat duct ruptures either within the epidermis leading to an intra-epidermal vesicle or at the epidermal dermal dermal junction leading to a subepidermal vesicle (O Brien, Sulzbergei, Jimmerman and Emerson). On the other hand, in mild cases the sweat duct does not break, as a rule, but sweat merely escapes from the sweat duct into the epidermis leading to the formation of an intra-epidermal "spongiotic" vesicle (Sulzberger and Zimmerman).

POMPHOLYX (DYSHIDROTIC ERUPTION)

This is a recurrent eruption of numerous deep seated vesicles occurring singly and in groups on the palms and the soles [Inflam]

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Lapı unl gaire et de la dermatite polymorphe de Duhring Brocq. Arch. belges dermat et syph 1 216 1939 Erythema Multiforme

Costello, M. J. Erythema multiforme exudativum. J. Invest. Dermat. 8 127, 1947. Winer L. H., and Lipschultz C E Comparative study of histology and cytology in vesiculating eruptions Arch Dermat & Syph 65 270, 1952

Burn

Moritz A R Studies of thermal injury III The pathology and pathogenesis of cutaneous burns Am | Path 23 915 1947

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Noninfectious Inflammatory Diseases

HRTICARIA

Urticaria is characterized by the presence of transient edematous than less or wheals, and is accompanied by considerable litching

Histopathology. An urticarial wheal shows edema, particularly of the upper dermis. The collagen bundles as well as the individual fibers are separated by edema. The collagenous substance appears swollen and stains poorly. It early wheals, i.e., those only a fen, min utes old, one finds either no inflammatory reaction or merely a slight perivascular infiltrate composed of lymphocytes. Wheals an



Fig. 39 Prurigo nodularis. There are hyperketations and co.

Hour old or older show a mild or even moderately severe lymphocytic-infiltrate around the capillaries (Torok and Lehner)

PRURIGO NODULARIS

chiefly

Histopathology One observes pronounced hyperkeratosis and acanthosis. There may be deep any aginations of the horny layer

ERYTHEMA NODOSUM

The lessons consist of tender red or haid red nodes which are slightly raised above the level of the skin. They vary from 1 to 5 cm in diameter and usually are limited to the anterior surfaces of the legs they may however occur elsewhere They involute within a few weeks without breaking down

Histopathology The histologic changes are located mainly in the upper portion of the subcutaneous rusue. The dermis merely shows a moderate amount of periosecular infiltrate composed predomi

nantly of lymphocytes

Invertigations one observes in the upper portion of the subcuta neous tissue a scattered infiltrate consisting mainly of neutrophils and Implicates Some-distrocytes and occasionally ensuring mainty of memorial and Implicates Some-distrocytes and occasionally ensuring the present flust plasma cells are absent. The inflammatory infiltrate extends as the disease progresses both upward toward the fatty tissue of the plant of the control of

irregular gutline NXabscess formation oxinecrosis occur

The blood vessels especially the vents may show severe involve ment so that many authors believe that a vascultus represents the primary and ptedominant lesion (Rotnes Grzybonski) Other authors however state that the blood sessels are not necessarily affected severely and may show only mild involvement (Pautsier and Woringer Lofgren and Wahlgren) In cases with vascular changes one observes especially in the larger veins invasion of the vascular 96

walls by the infimmatory infiltrate and marked endothelial proliferation (Fig. 40). However, complete occlusion and thrombosis are

Epithelioid and giant cells are absent in early lesions. However occasionally one finds small nodules composed of histocytes lying either in radial arrangement or in palisade like arrangement around

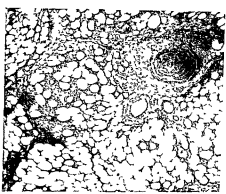


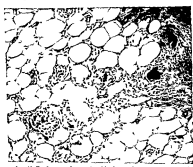
Fig. 40 Erythema nodosum A nonspecific inflammatory infiltrate extends in small scattered patches through the subcutaneous fat. A large subcutaneous vein shows endothelial proliferation and invasion of its wall by inflammatory cells (%50)

a small central fissure (Fig. 41) (Misscher). Not infrequently these nodules are permeated by neutrophils. Since they are found in no other disease, they are of considerable diagnostic value (Nubé).

Older lesions show lewer neutrophils and more lymphocytes than young lesions. Often giant cells are present at this stage. The giant cells usually of the foreign body type may be found outside of or within foci of epithelioid cells. In the latter case, the arrangement simulates that found in tuberculosis (Rotnes). However, casellion is always absent.

Differential Diagnosis For differentiation from crythema induratum see page 186 In, cases of crythema nodosum showing severe vascular involvement, periarterius nodosa must be excluded. In the

latter disease however the autenest are predominantly involved one observes necrosis of vascular walls and the infiltrate usually contains a large percentage of cosmopfuls odular vasculitis greatly resembles the late stage of erythema nodosum It differs from erythema nodosum by showing a larger degree of vascular involvement including in



to 41 Erythema nodosum. An older lesion shows two small nodules composed of histocytes and to the right several giant cells of the for eigh body type (×200).

solvement of vessels of large caliber. However, it is possible that it represents a variant of erythema nodosum.

NODULAR VASCULITIS

Clinically this disease is characterized by slightly painful nodules occurring chiefly on the legs. They generally neeser for a

receives (Fig. 12) In addition one observes a various of ...

walls by the inflammatory infiltrate and marked endothelial proliferation (Fig. 40). However, complete occlusion and thrombosis are

Epithelioid and giant cells are absent in early lesions. However, occasionally one finds small nodules composed of histocytes lying either in radial arrangement, or in palisade like arrangement around

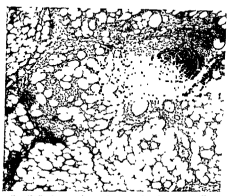


Fig 40 Erythema nodosum, A nonspecific inflammatory infiltrate extends in small, scattered patches through the subcutaneous fat A large subcutaneous vein shows endothelial proliferation and invasion of its wall by inflammatory cells (×50)

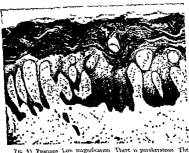
a small central fissure (Fig. 41) (Miescher) Not infrequently, these nodules are permeated by neutrophils. Since they are found in no other disease, they are of considerable diagnostic value (Nubé)

Blder lesions show fewer neutrophils and more lymphocytes than soung lesions. Often giant cells are present at this stage. The giant cells, usually of the foreign-body type, may be found outside of, or within foci of, epithelioid cells. In the latter case, the arrangement simulates that found in tuberculosis (Rotnes). However, cascation is always absent.

Differential Diagnosis. For differentiation from crythema induratum, see page 186 In, cases of erythema nodosum showing severe vascular involvement, persarteritis nodosa must be excluded. In the

ıs absent

The stratum palpighii is thinijed above the papillae, often to only two or three layers of cells. The Yete ridges show considerable clongation They often are slender in their upper portion and thickened in



rete ridges are clongated and thickened in their lower portion. There are edem and clubbing of the papillae (×50)

their lower portion. They may show branching at their bases, causing

may be found scattered through the stratum malpights

Corresponding to the elongation and the branching of the rete ridges the appliae are elongated and tortuous. The upper part of the papillae is edematous and club shaped. The capillaries in the papillae are dilated and torthons and show slight thickening of their walls These changes in the capillaties can be demonstrated best by the use of the Hotchkiss McManus stain (Stoughton and Wells). A mild to moderately severe inflammatory infiltrate is present in the upper dermis, particularly in the papillae It consists of lymphocyte

It is probable that nodular vasculitis is a variant of erythema nodosum. For their differential diagnosis see under erythema no dosum, see page 97.

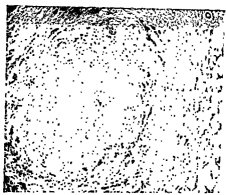


Fig. 42. Nodular vasculitis. The wall of a large subcutineous vein is greatly thickened and infiltrated with inflammatory cells. The lumen is obliterated (×50)

PSOR IASIS

Psoriasis is a chronic disorder characterized by dull red or brownish papules and plaques. The lesions are sharply demarcated, dry and usually covered with layers of fine silvery scales. As the scales are removed by gentle curettage, one frequently sees characteristic fine bleeding points. In severe cases, the disease may affect the entire skin and present the clinical picture of generalized exfoliative dermatitis (exfoliative psoriasis).

Histopathology. Histologically, psoriasis is characterized by (I) parakeratosis, (2) thinning of the suprapapillary portions of the stratum malpighii, (3) elongation of the rete ridges, (4) edema and clubbing of the papillae and (5) Munro micro-abscesses (Fig. 43)

Corresponding to the layered silvery scaling observed clinically, the horny layer is considerably thickened and consists predominantly of parakeratotic cells, arranged in lamellae with air spaces in between.

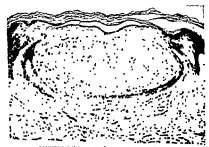
edema of the rete cells overlying the papillae and (4) capillary dilatuion in the tips of the papillae

EXPOLATIVE PSORARS The histologic picture of exfoliative psoria six may still show enough of the characteristics of psoriasis to allow this diagnosis Frequently however the appearance is indistinguish able from that of exfoliative dermatus due to other causes

Differential Disgnosis The histologic picture of psoriasis may re semble that of tichen simplex chropicus. For their differentiation, see prige 73. Differentiation, from borrheic dermatitis is not always possible. Schottheic dermatitis may show all the features of psoriasis, though less pronounced than in psoriasis. In addition, however, one finds a fair degree of spongosis, which in psoriasis is either very slight or obsent.

PUSTULOSIS PALMARIS ET PI ANTARIS (PUSTULAR PSORIASIS PUSTULAR BACTERID)

Two diseases have been described in the literature under the name of pustular psoriasis generalized pustular psoriasis and pustular



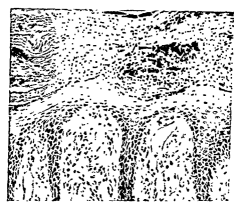
ntra-epidermal unilocular pustule containing many neutrophils is present

portrais of the paims and the soles. Neither of them however, is form of psoriasis. Actually, pusually lesions do not occur in psoriasis. So-called generalized pusualar psoriasis is identical with the gen

100

and histiocytes, except in early lesions which, in addition, show polymorphonuclear lenkocytes in the infiltrated Plasma cells are found only rargly and posinophils hardly ever

The Munro micro abscesses (Fig. 44) are located either in the stratum corneum or directly beneath it. They represent small ac-



Fit 44 Psorpasis High magnification of Figure 43 \ \text{Munto micro} abscess (M) is located within the parakeratotic born) hyer. The supripipillary portions of the stratum malpiglin are thanned and show intracellular edema. Intercellular edema, however is absent. The cipillaries in the tips of the papillac are dilated (X200)

cumulations of neutrophils which have migrated there through the epidern's Munro micro abscesses, as a rule, are found easily infearly lesions. Includer lesions, they are few in number or even absent. Thus, their absence does not rule out psoriasis. Neither does their presence establish a diagnosis of psoriasis, since Munro micro abscesses may occur also in sebornien dermatitis, accidernatitis con timus (Hallopeaus keratosis blennorrhagica and parapsoriasis guitata.

The fleeding points which may be produced by gentle scraping of the skin correspond to the apices of papillae They are due to the following histologic changes (1) parakeratosis, (2) thinning of the stratum malpighia above the ups of the papillae, (3) intracellular

walls thus form a sponge like network in the interstices of which neutrophils continue to accumultue (Fig. 46). As the pustule ages the cellular walls gradually break in the center of the pustule so that



he 46 Acrodemantin commun of Hallopeau There is acanthous with elongation of the zete radger. The upper stratum malpighu contains a spongiform possible the cellular walls of the elemantous squareous cells fram a spongs the twork in the interstuces of which resecrophils have accumulated (x500)

a large cavity forms. It is e periphers of the pustule however, the network persists for a much longer time.

Differential Diagnosis. The spongulorin pustule of kogoj represents a striking histologic lesion and is characteristic of accoderma titus continua. However, it is not diagnostic of that disease since it occurs also in three other diseases, imperigo herpetiformis keratous blennorthagica and Reiter's disease. Differentiation of these four eralized form of acrodermatitis continua (Lapiere, Bruck) and will

be described there (see below)

Pustular psoriasts of the prims and the soles, a term introduced by Barber, is a misnomer. This disease is now referred to frequently as either pustular bacterid (Andrews) or acrodermatitis pustuloss (Sachs, MacKee and Rothstein). The term pustulosis primaris et plantaris is preferable, however.

Pustulosis pulmaris explantaris is a chronic, indolent disorder limited to the palms and the soles and characterized by the appear ance of crops of deep seated pustules within areas of erythema and

scaling

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Histopathology. The histologic picture does not resemble that of psoriasis. A large, intra epidermal, unifocular pustule is present (Fig 45). It contains many neutrophils and disintegrated cellular elements. The epidermis surrounding the pustule shows slightly canthosis with little or no spongiosis. In the dermis underlying the pustule, a mod crately severe inflammatory infiltrate is present, composed mainly of lymphocytes and histocytes but containing also a few neutrophils. Histologic features of isoriasis, such as parakeratosis elongation of the reterridges. Minning of the suprappillary portions of the stratum malpighu and dilatation of the Vapillary capillaries are always absent (Sachs and Scannone, Sachs, MacKee and Rotistem)

ACRODERMATITIS CONTINUA

(Halloperu)

Acrodermatitis continua of Hallopeau, a chronic disease hiving pustules as a primity lesion, usually is limited to the hands and the feet. The distal portions of the fingers and the toes are predominantly involved. Occasionally, however, the disease is generalized. The affected areas are dirk red, dry, shiny and scaling and are studded with shallow pustules. Lesions located on the distal portions of the fingers and the toes may cause atrophy of the skin and loss of nails.

Histopathology. The epiderinis shows parakeratosis and moderate acanthosis with elongition of the rete ridges. An inflammator in filtrate containing many neutrophils but few or no cosmophils is present in the dermis. Many neutrophils are seen invading the

epidermis

However, the characteristic lesion consists of the so called spongi form pustule of Kogoj. This type of pustule forms in the uppermost stratum malpighii through the migration of neutrophils into edema tous squamous cells. This invasion cruses disintegration of the cyto plasm and of the nucleus but not of the cellular walls. The cellular

walls thus form a sponge like network in the interstices of which neutrophils continue to accumulate (Fig. 46). As the pustule 'gges, the cellular walls gradually break in the center of the pustule so that



his 6. Accodematius continues of Hallopean There is acanthons with elongation of the rete ridges. The upper action in Judgella contains a spongiform pustule the reliable walk of the edematous squamous cells form a sponge like net work in the intersities of which neutrophils have accumulated (x500).

a large cavity forms. At the periphery of the pustule however, the network persists for a much longer time

Differential Diagnosis The spongiform quistule of kogoj represents a striking histologic lesion and is characteristic of acroderma titis continua. However, it is not diagnosic of that disease since it occurs also in three other diseases, impetigo herpetiformis, keratosis blennorthagica and Rener's disease. Differentiation of these four 104

diseases may be impossible. However, in impetigo herpetiformis, the infiltrate contains, as a rule, a large number of eosinophils (see below). Older lesions of keratosis blemorrhagica and Reiter's dis ease often can be differentiated from acrodermatitis continua by the presence of a greatly thickened horny layer (see page 172)

IMPETIGO HERPETIFORMIS

This rare, usually fatal, disease is seen mainly in pregnant women and occasionally in hypopirathyroidism. It is characterized by the presence of pustules set in groups or in circinate arrangement on erythematous patches

Histopathology. The histologic picture, like that of acrodermatitis continua, is characterized by the presence of the spongiform pusule (Kogoj) (See "Acrodermatitis Continua" for a description of the spongiform pustule) The dermal infiltrate as well as the pustule

contain a conspicuous number of cosmophils (Rost)

Differential Diagnosis. The presence of many cosmophils aids in the differentiation of impetigo herpetiformis from the other three diseases with spongiform pustules acrodermatitis continua, kera tosis blennorrhagica and Reiter's disease. The latter two diseases, in addition, often show conspicuous by perkeratosis in their older legions

PARAPSORIASIS

Paraprortasis comprises a group of rare dermatoses. Three of the four forms of parapsoriasis are characterized by an asymptomatic maculopapular eruption of slow evolution and marked chromicity. These forms are parapsortasis guttata, parapsortasis lichenoides (parakeratosis variegata) and parapsortasis en plaques. In parapsorta sis guttata, one observes especially on the trunk numerous papules which may or may not be covered with fine scales. In parapsoriasis ment In parapsortasis emplaques, well defined parties and plumes of various size and shape occur. The fourth form of parapsortasis parapsoriasis, carioliformis of Habermann, ilso called pityriasis lichenoides et varioliformis acuta differs from the previously men the through the state of the st and papulonecrotic lesions terminating in varioliform scars

Histopathology. None of the four forms of prraportasis shows a diagnostic histologic picture Varapsoriasis guitata, prrapsoriasis ichenoides and parapsoriasis en pluques show the histologic picture of chronic dermatius, while parapsoriasis Varioliformis shows an acute inflammatory process with foci of necrosis

the infiltrate in the manner of a claw clutching a hall

The duse of the disease is not known Ellis and Hill regard lichen mitidus as closely related to liciten planus because they found that the two diseases may occur together and histologic sections obtained from patients with lichen planus may show lichen mitidus like lessons

LICHEN STRIATUS

Lichen stratus is an uncommon eruption which as a rule occurs in children It manifests uself usually on the extremities as a long band composed of <u>small lichenoid papilles</u>. The eruption appears suddenly and involutes within a few weeks or months. Itching is absent

Histopathology The histologic changes are similar to those of neurodermatitis circumscripta. In some cases the chronic inflammatory inflittate is more markedly pertuascular than in neurodermatitis circumscripta and surrounds the vessels of the upper dermis 18 densely packed manites (Senear and Caro Pinkus).

PITYRIASIS RUBRA PILARIS

The primary lesions are reddish follicular papules. They gradually coale of the Ultimately most of the coale of the coale

of the fingers

Histopathology The essential pathologic process is follicular by practicular in addition there is diffuse hyperkerators with spotted parakerators. The epidermis shows irregular acanthosis usually of mild degree There often is liquefaction degeneration of the basic cells (Brunsting and Sheard) In the upper dermis a mild chronic inflammatory infiltrate is observed around the blood vessels.

Differential Diagnosis For a differential diagnosis from phrynoderma (vitamin A deficiency) see page 286

GRANGLOMA FACIALE (EOSINOPHILIC GRANGLOMA OF THE FACE)

This disorder only recently established as a disease entity consists of soft purplish slowly growing and asymptomatic patches limited to the face Except for prominence of the follicular openings the surface of the skin appears normal

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Histopathology. A dense, granulomatous infiltrate is located mainly in the upper half of the dermis (Fig. 52) Quite characteristically, the infiltrate does not invade the epidermis or the pilosebacous/ppendages but is separated from them by a narrow zone of normal collagen. The pilosebacous appendages are well preserved. Two stages can be differentiated in the development of the

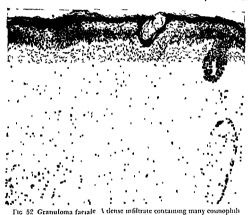
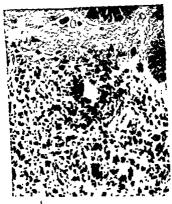


Fig. 52 Granuloma faciale \(^1\) dense infiltrate containing many cosmophils is present in the dermis. The infiltrate does not invade the epidermis or the sebaccous gland shown on the right but is separated from them by a zone of normal collagen (\times^{100})

infiltrate, an early 'leukocytic and a late "fibrotic" stage The two stages may occur together in the samp lesion

In the darly "leukocytic" stage, leosinophils predominate, but neutrophils and histocytes are found in large number and, in ad dition, a few lymphocytes, plasma cells and mast cells. The infiltrate thus has a polymorphous appearance. The capillaries are dilited and surrounded by degenerated collagen (Fig. 53). In some cases con siderable amounts of hemosiderin are present in the upper dermis within and outside of histocytes (Lever, Lane, Downing and Spingler). The presence of foam cells has been described in one case (Lever).

In the Me 'fibrotic' stage, the infiltrate is broken up into variously sized patches by strands of collagen The number of cosinophils and neutrophils is greatly reduced, and plasma cells' hymphocytes and fibroblasts predominate The capillaries often have fibrotic walls



of cells cosmophils and histocytes predominate (x400)

Differential Diagnosis. The arrangement and the composition of the infiltrate in granuloma faciale result in a diagnostic histologic picture, with no resemblance to that of distinguishing granuloma with which the disease was at first confused (Lever) Although cosmophilic grinulomi which represents an abortive form of Hand Schuller Christian disease (see page 265) also contains many cosmophilis and histocites us histologic appearance differs from that of granuloma faciale by the following

larger and he in patch the epidermis (Lever and Leeper) For a differentiation from ery thema elevatum diutinum, see below

TRYTHEMA ELEVATUM DIUTINUM

This rare disorder shows persistent, jed to purple nodules with some tendency to grouping about the joints, especially on the dorsa of the <u>hands</u>, the wisits and the <u>elboys</u>. The nodules at first are soft but later, due to fibrous transformation, become hard

Histopathology. Frythema elevatum diutinum has a fairly drig nostic histologic picture which was first described by Weidman and Besincon, and then by Ketron and by Weiss as well as by others. The disease described in 1891 by Crocker and Williams as crythema elevatum diutinum of the Bury type is now widely regarded as a variant or late stage of granulomy annulary. (Haber and Russell)

Frythema elevatum dintimum, in its early strage, shows a dense, predominantly perivascular infiltrate composed largely of neutro phils intermingled with some lymphocytes and histocytes. In most cases, a peculiar hyaline degeneration of the reticulum fibers around the capillaries has been observed. This degeneration has been referred to by Weidman and Besancon and by Ketron as formation of toxic hyalin." This toxic hyalin has a smooth, glassy appearance and stains phensely pink with equin

In the Atte, fibrous stage, the cellular infiltrate is much less pro nounced and extensive fibrosis is present. The capillaries may still show their mantle of toxic by alm or may merely show fibrous thickening (Ketron)

Differential Diagnosis The histologic appearance of erythema

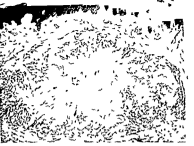
a zone of normal collagen beneath the epiderinis and the presence of coxic hyalin—about the capillaries

GRANULOMA ANNULARE

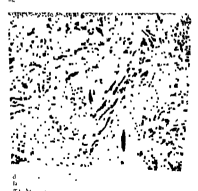
The lesions which are found most commonly on the hands and the feet, consist of small, firm, pale red nodules which tend to be ngement. The disease is chronic

anuloma annulare is character

ived by focal degeneration of the collagen in the dermis deposits of mucin between the degenerated collagenous bundles and rective inflammation and fibrosis. The degeneration of collagen within the foci may be complete or incomplete. Some cases show foci of both complete and incomplete degeneration and others show only one



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large focus of complete colligen degeneration. In the majority of cases however one finds multiple foct of incomplete degeneration without any fact of complete degeneration.

In foci of complete degeneration one finds a shaiply demarcated area of congulation necrosis surrounded by an infiltrate of histocytes fibroblasts and lymphocytes in a radial arrangement (Lig 54)

In foci of momplete degeneration one finds ill defined areas in which collagen fibers are found in various stages of degeneration ranging from mild homogenization and fragmentation to granular degeneration and congulation necrosis. A considerable amount of the collagen in such areas may appear normal. Lymphocytes histocytes and fibroblasts infiltrate between the patrially degenerated collagen fibers and new collagen is being laid down. Thus the affected areas of the dermis present a completely disorderly arrangement of the collagen bundles (Fig. 55).

In both types of collegen degeneration though more frequently in areas of incomplete degeneration fine threads and granules of mucin are deposited between the degenerated collegen bundles. The mucin string light blue with routine string and red when Best's mucicarmine string is used. (See Plate 1.)

In trens of collingen degeneration, the chistic usine shows fragmentation and may be largely destroyed. In some areas, there may even be greater destruction of elastic than of connective usine. A moderate or even considerable amount of Competitive usine is moderate or even considerable amount of Completive usine in Intervals of the blood vessels outside the areas of degeneration. The walls of the blood vessels show no pathologic changes however except occasional mild endothelial proliferation. In some cases, a few grant cells of the foreign body type are present. They usually at situated near the periphery of the infiltrate and are not associated with any zone of necrosis (Prunty and Montgomery).

Differential Diagnosis. The type of granuloma annulare showing multiple foct of incomplete collagen degeneration may greatly resemble necrobiosis lipoidica which shows this stone pattern of collagen degeneration (see page 269). However, acrobiosis lipoidica differs from granuloma annulare by the presence of viscular clanges of lipid material and of many foreign body grant cells and by the absence of much

The type of granuloma annulare showing one or several large foci of complete collagen degeneration any be almost indistinguish able from the subcutaneous nodes of meumatoid arthritis (see below). However, in the nodes of rheumatoid arthritis the areas of degen the dermis

red cellular

eaction, mainly histocytes and fibroblasts, while in granuloma an iulare there are, in addition, many lymphocytes (Bowers)

SUBCUTANEOUS NODULES OF RHEUMATIC FEVER AND RHEUMATOID ARTHRITIS

In both rheumatic fever and rheumatoid arthritis, small firm modules may form in the subcutaneous tissue. The commonest sites

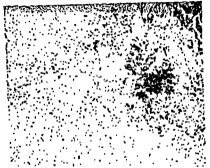


Fig. 56 Subcutaneous nodule of theumatoid arithmin. There is a large central rate of necrous surrounded by historytes in palisade arrangement (×100).

are about the elbows, the knees and the ankles. Their size varies from a few millimeters to 2 cm

Histopathology The subcutaneous nodules found in these two diseases are similar in ming of their pathologic features. In both diseases they are evaposed of three reasonably well-defined zones. (1) a central zone, or zone of necrous, (2) an intermediate zone, comprising proliferating histopaciae cells and (3) a pertipheral zone, consisting of chrone influentatory cells (fementi, Teller and Bauer). However, exidative changes presail in the nodules of theumatic fever, Afolderative and necrotic changes in those of pheumatical arthritis.

large focus of complete collagen degeneration. In the majority of cases, however, one finds multiple foci of incomplete degeneration without any foci of complete degeneration.

In foci of complete degeneration, one finds a sharply demarched area of coagulation necrosis surrounded by an infiltrate of histocytes, fibroblasts and lymphocytes in a radial arrangement (Fig. 54)

In foci of meaning the degeneration, one finds ill defined areas in which collagen fibers are found in various stages of degeneration ranging from mild homogenization and fragmentation to granular degeneration and coagulation necrosis. A considerable amount of the collagen in such areas may appear normal Lymphocytes, histocytes and fibroblasts infiltrate between the partially degenerated collagen fibers and new collagen is being laid down. Thus, the affected areas of the dermis present a completely disorderly arrangement of the collagen bundles (Fig. 55).

In both types of collagen degeneration, though more frequently in areas of incomplete degeneration, fine threads and granules of mucin are deposited between the degenerated collagen bundles. The mucin stains light blue with routine stains and red when Best's mucicarmine stain is used (See Plate 1)

In areas of collagen degeneration, the ensure ussue shows fragmentation and may be largely destroyed. In some areas, there may even be greater destruction of elastic than of connective tissue A moderate or even considerable amount of ymphocytic infiltration is found around the <a href="https://linearchy.org

Differential Diagnosis. The type of granuloma annulare showing multiple loci of incomplete collagen degeneration may greatly resemble necrobiosis lipoidica which shows this stone pattern of collagen degeneration (see page 269). However, recrobiosis lipoidica differs from granuloma annulare by the presence of viscular changes, of lipid material and of many foreign body grant cells and by the absence of much

absence or interest. The type of granuloma annulare showing one or several large foci of complete collagen degeneration may be almost indistinguish able from the subcutaneous nodes of encumatoid arthritis (see below). However, in the nodes of rheumatoid arthritis the areas of degeneration are usually larger, they are more deeply located in the dermis or even in the subcuttis, and they are encircled by a less varied cellular

After many years' duration, an atrophic stage may be reached in which the findings are no longer diagnostic, since one sees merely an atrophic epidermis and an atrophic, fibrotic dermis without inflammatory infiltrate.



Fig. 57. Acrodermantis chronica atrophicans. There is actrophy of the stratum malpighii. A handlide infiltrate is separated from the epidermis by a narrow rome of normal collagen. The dermis shows intensitial edema and atrophy of the collagen bundles. Because of this arophy, the thickness of the dermis is markedly decreased and the sweat glands lie unusually close to the pedermis [X100].

The bindlike areas of fibrosis resembling scleroderma and the

able from that of scleroderma

RADIODERMATITIS

An early (acute) and late (chronic) stage of radiodermatitis are recognized. The early stage occurs within a few days after adminis

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In rheumatic fever, the central zone shows "fibrinoid degeneration" (For a discussion of fibrinoid degeneration, see "Acute Systemic Lupus Erythematosus," page 297.) The intermediate zone is composed of edematous collagen infiltrated with inflammatory cells, including many large mononuclear cells (histocytes) which resemble the predominant cells of myocardial Aschoff nodules.

In heumatoid arthritis, the central zone shows necrosis of all pre existing collagen, and the intermediate zone distinct palisading of histocytes and fibroblasts around the central zone of necrosis (Fig.

56)

Differential Diagnosis. For differentiation of the nodules of rheumatoid arthritis from granuloma annulare, see 'Granuloma Annulare,' page 116

ACRODERMATITIS CHRONICA ATROPHICANS

This condition affects the extremities. The skin presents a bluish red or a brownish, atrophic, wrinkled appearance. Fine scaling is usually present. Because of decrease in the amount of subcutaneous fat, the subcutaneous veins are clearly visible. In some cases, bandlike areas of fibrosis, resembling scleroderma, and fibrous nodules are observed, especially along the ulna as "ulnar bands" and along the table.

Histopathology. The histologic picture is characteristic. One ob serves mild to moderate hyperkeratosis and atrophy of the epidermis, with absence of the rete ridges. Just beneath the epidermis, there is a narrow zone of connective ussue separating a dense band of in flammatory infiltrate from the epidermis. In addition to this band like infiltrate, one finds scattered areas of inflammatory infiltration throughout the dermis, particularly around the blood vessels. The infiltrate is composed predominantly of lymphocytes, but also contains histocytes. Chromatophores laden with melanin or hemosiderin may be present. The entire dermis shows interstitual edema and atrophy of the collagenous bundles. The atrophy results in a gradual decrease of the dermis to a half or a quarter of its normal thickness (Fig. 57).

The sebaceous glands and the hairs undergo atrophy early in the disease and usually are entirely absent. However, the sweat glands are, as a rule, preserved. Because of the thinness of the dermis, they he unusually close to the epidermis. The blood vessels are dilated and may show endothelial proliferation.

The subcutaneous tissue shows decided atrophy. The fat cells vary in size and are irregular in shape. Foci of inflammatory infiltration

and a varying degree of fibrosis are present

changes An inflammatory infiltrate is seen throughout the dermis, especially about the appendages. The blood vessels are dilated and reteal ederm of their walls and endothelial proliferation. The collegen bundles show edems and homogenization. In severe cases, the



Fig. 59 Late radiodermatitis. The vessels (V) show

epidernus and the upper dermis undergo necrosis. The area of necrosis is invaded and surrounded by polymorphonuclear leukocytes

Histopathology of Late (Chronic) Radiodermatitis, The epiderms is irregular showing atrophy in some areas and acanthosis with hyper keratosis in others. The rete cells frequently show atypicality, such as disorderly arrangement, individual cell dyskeratosis and an increased number of minotic figures. Thus the changes may resemble those of Bowens disease (intra-epidermal squamous-cell carenoma). In addition there often is irregular downward growth of the epidermis. The

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tration of a massive dose of roentgen rays or radium. At first, there is erythema, which is followed by desquamation and pigmentation. If the dose administered is sufficiently large, ulceration will ensue. The late (chronic) stage of radiodermatitis occurs from a few months.



Fit 58 Late radiodermatitis The epidermis shows acm thous and downward grouth around a telangecertic blood tested Many irregularly diluted lymphatics are located directly beneath the epidermis. The collagen shows degeneration (£/100)

to several years after the administration of large amounts of roenigen rays or radium. The skin shows atrophy with interspersed areas of hyperkeratosis, irregular hyperpigmentation, telangiectases and loss of hair. Ulceration may be present. The hyperkeratoses may develop into squamous cell carcinoma.

Histopathology of Early (Acute) Radiodermatitis. The cells of the epidermis, particularly those of the basal layer, are hydropic and their nuclei show pyknosis. The epithelial cells of the hair follicles, the sebaceous glands and the sweat glands may show similar degenerative.

Histopathology The histologic appearance usually is that of a nonspecific chronic derinaturis (see page 70). However, the erythemotous lessons frequently present areas of liquefaction degeneration of the basal layer, and edema together with a patchy predominantly symphocytic infiltrate in the upper dermis. Differentiation from lupus erythematosis may then be impossible. However, in some cases of solar dermaturis a moderate number of cosinophils is present in the infiltrate. Since eosinophils are generally, not observed in lupus erythematosis their presence aids greatly in the differentiation of the two diseases (Lamb).

HIDROCASTOMA

This condition is characterized by the presence on the face of numerous clear tense deep-seated vesicles varying in size from a few millimeters to 1 continuer

Histopathology On histologic examination one observes in the dermis large cystic cauties lined by epithelial cells which usually are arranged in two layers. Small papillary projections into the lumina of the cysts have been described (hened) and Lehner).

Most authors regard the cysts as caused by retention of sweat in the dermit portion of eccrine sweat discis. However, kenedy and Lehner believe that the cysts are not merely she result of retention of sweat and passive distation of the exerctory ducts but are due to a pro-liferation of the exerctory duct epithelium, which leads to occlusion of the ducts.

ACNL VULGARIS

Acne sulgarus a disease occurring predominantly in pubers; affects the face the upper chest and the upper back. The primary lesson is the comedo As a foreign body reaction to the comedo and due to secondary infection follicular pipules and pustitles and subcuta nosus abserses may occur.

Histopathology Histologically acne vulgitis tepresents a perifollicultus occurring around a comedo. Comedones are composed of sebum and keratoric as well as parakeratoric cells and are located in the pilosebaccous follicles.

In the early acne lesson one finds an infiltrate of tymphocytes and family acress the control of the property of the property of the control of the control of the control of the fragmentation of sebaceous glands. The atrophy and the fragmentation of sebaceous glands probably are due to the stass and the pressure caused by the contedo within the follicle. The fragmentation of sebaceous glands in turn causes an inflammatory reaction (Lynch). Occasionally foreign body giant cells are found in

rete ridges may grow around telangiectatic vessels, which may thus become completely enclosed in the epidermis (Fig. 58).

In the dermis, the collagenous fibers are swollen and sclerotic In some areas, they are broken apart and stain basophilic Formation of new, young connective tissue is found throughout the dermis, but especially around the blood vessels Directly beneath the epidermis, one often sees numerous irregularly dilated lymphatics as well as lymphedema

However, the fundamental pathologic change is that of fibrotic thickening of the walls of the vessels in the deeper portions of the dermis, leading to occlusive changes in the lumina (Fig. 59) Some of the vessels show thrombosis and recanalization. The vessels nearest

to the epidermis may show telangiectasia

Hair follicles and sebaceous glands are absent, but the sweat glands usually are preserved, except in areas of third degree injury. The elastic tissue is, as a rule, less damaged than the collagenous tissue—at least in milder degrees of radiodermatus. In severe reactions, it is destroyed. With Foot's reticulum stain, the reticulum fibers are seen to be increased throughout the dermis wherever there is formation of young connective tissue, but especially around blood vessels.

In severe cases of chronic radiodermatitis, ulceration occurs. The deep lying, large blood vessels beneath the ulcers, as a rule, show

complete obliteration

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The above described epidermal changes of dyskeratosis and intra epidermal carcinoma frequently lead on to invasive squamous cell carcinoma. A fairly large percentage of them are of high (grade IV) malignancy and of the spindle celled variety (see page 333). In rare instances, basal cell epitheliomas develop in areas of radiodermatitis (Anderson and Anderson) (see under Basal Cell Epithelioma," page 370). The occurrence of sarcoma, on the other hand is not fully established. Such instances have been reported (Blom Ides, Gentele), but always with the reservation that the diagnosis of spindle celled squamous-cell carcinoma could not be ruled out conclusively because this differentiation may be nearly impossible (see page 410). A point in favor of their being carcinoma is the fact that they are, in contrast to fibrosarcoma, radiosensitive (Blom Ides).

SOLAR DERMATITIS

Solar dermutitis may occur as erythematous, urticarial, papular or vesicular lesions in areas exposed to the sun. The erythematous lesions may resemble lupus erythematosus in their clinical appearance by showing a livid red color and slight induration.

but he concedes that differentiation of the two diseases by histologic means may be impossible. Additional but not obligatory findings in the papular type of rosacea are superficial perifoliculitis. with intrafollicular abscess formation (pustules) and dilatation of capil laries. There is no hypertrophy of the schaceous glands.

In the glandular hypertupus of the schools glands are in creased in size and number. The orifices of the sebaceous glands are in creased in size and number. The orifices of the sebaceous glands are dilated and filled with ketatin and sebum. There is an associated hypertrophy of the connectine tissue. The blood vessels are dilated a thronic inflammatory infiltrite is present around the vessels. Perfolliculties with intrafollocular absects formation may be observed.

FON FORDICE DISEASE

This disease occurs only in women It is characterized by an erup-

spectine glands is the limitation of the eruption to the areas where spectine glands occur and the onset of the disease at puberty when the apocrine glands begin to function. Shelley and Hurley have expressed the opition that Fox Fordyce disease may be basically a miliaria apocrina—in other words an apocrine sweat recention miliaria.

Histopathology Histologic examination reveals acanthosis with hyperkeratosis. The reteridges are elongated \text{\chi} moderate degree of round cell infiltration is present around the blood vessels in the upper dermis. The appearance thus is that of a chronic dermatitis.

The apocrine glands appert normal (Royburgh) The statement frequently found in the literature ('slifen) that in Foy Fordyce dis ease the sweat glands are dilated is due to the fact that many authors have misinterpreted the apocrine glands as dilated sweat glands.

ALOPECIA AREATA

Mopecia reats is characterized by loss of hair in one or

involved (alopecta totalis) in which case the loss of hair is usually permanent

Histopathology The hard follicles and the hard bulbs are greatly reduced in size. The hard bulbs instead of being deep in the subcutts are located quite shiph, up in the definits. The hair follicles contain foose keratin and no hair. In long standing cases there is

the infiltrate but there is no evidence of tuberculoid reaction. At a later stage, histocytes and fibroblasts appear and fibrosis occurs

In lessons in which secondary betterial infection is present one finds disintegration of the follicle and abscess formation. Foreign body grant cells are then frequently found around the remnants of the follicle.

ACNE VARIOLIFORMIS (ACNE NECROTICA MILIARIS)

Acue varioliformis is characterized by the presence of small indo lent papules and pustules along the frontal hardine. The lesions undergo central necrosis and tend to heal with pitted sears.

Acte necrotics militaris of the scalp is a diminutive variant of acte varioliformis. Because of the superficial location of the necrosis no hair loss ensues

Histopathology The histologic changes of acide necrotica are similar to those of acide vulgaris except that the penfolliculitis is more limited in extent and almost invariably results in central necrosis Blood vessels adjacent to the necrotic area may be thrombosed. Healing takes place with fibrosis and sear formation.

Acte pecrotics militris of the scalp shows the same histologic picture as that of actic varioblogmis (Montgomery)

ACNT ROSACTA

Acne rosacea occurs in patients with seborrher and affects the central portion of the face. Two forms of rosacea exist the papular form and the glandular hyperplastic form. The two forms may occur to gether. The papular form is characterized by erythema papules pustules and telangiectases. The glandular hyperplastic form causes enlargement of the nose called rhipophyma.

Histopathology In the papular variety the infiltrate is either diffusely distributed throughout the dermis or arranged about the hur follicles and the sebaceous glands. As a rule it is banal in character and composed largely of lymphocytes together with some histocytes and plasma cells. However, a significant number of patients show in addition foci of epithelioid cells and Langhans grant cells occasionally, lying in true tubercle formation, so that the histologic picture is indistinguishable from that of cutaneous tuberculosis. Miescher who found tuberculoid for im 80 per cent of 39 patients with acine rosacca concluded that the rosacealise tuberculoid Lewandowsky and the papular type of tene rosacca represent the same disease (see page 183). On the other hand, Laymon who observed tuberculoid structures in only 11 per cent of 138 patients with acine rosacca does not think that the two diseases are identical

Purpura occurs as the result of either noninflammatory or inflam matory changes in the walls of small blood vessels

NONINFLAMMATORY PURPURA

The following types of purpura are noninflammatory stasss purpura due to increased intracapillary pressure senile purpura due to degeneration of the dermal collagen in exposed areas of the skin scurry due to reduction of the intercellular material between endo thehal cells (thrombocytopenic purpura due to inadequate formation of blood platelets and febrile purpura with platelet throm boss a rare systemic fatal disease

Histopathology In staiss purpura scurvy and thrombocytopenic purpura no visible vascular changes are present and the only ab normality consists of the presence of extravasated red blood cells and at a later stage of hemosiderin. In senile purpura one observes in addition severe degenerative changes of the dermal collagen as seen in senile elastosis (see page 157) (Tattersall and Seville). In febrile purpura with platelet thrombosis—one observes in the capitlaries of the skin as well as of many internal organs degeneration and pro-tiferation of endothelril cells and obstructing thrombi composed of platelets (Trobaugh).

INFLAMMATORY PURPURA (VASCULITIS)

The following types of purpura are caused by inflammatory changes in the walls of small blood vessels (asculiets) bacterial purpura due to meningococi. Streptococcus viridans or other microorganisms anaphylactoid or Schoenlein Henoch purpura which is of unknown genesis and often associated with joint pains gastrometismal bleeding and occasionally with glomerulonephritis purpura due to drug allergy and purpura associated with periarterius nodosa.

Histopathology There are vascular thanges and an inflammatory infiltrate in addition to the extravastion of red blood cells

For the histologic changes in the purpuric lesions of meningococcenta and subactile bacterial endocarditis see pages 170 and 171 respectively. For purpura associated with periarteritis nodosa see page 511 Purpura due to drug allergy has the same histologic picture as that of anaphylactoid purpura

NATIONAL TOIN PURPLEA. Early lesions show prominent vascular changes in the upper dermis consisting of swelling and degeneration of endothelial cells and occasionally of focal recrois of the vascular wall (Gardner Levinson). A rather severe inflammatory infilirate is

gross thinning of the dermis, which may be reduced to half its normal thickness (Dillaha and Rothman). In cases of secent onset, a mild to moderately severe inflammatory infiltrate composed of symphocytes is seen in the deeper dermis about the vessels, the sebuceous glands and the hair follicles, while, in cases of more than 1 year's duration, the inflammatory reaction has subsided (Laymon). The sebuceous and the save a glands appear normal throughout the course of the disease.

ALOPECIA CICATRISATA (PSEUDOPELADE BROCO)

In alopecia cicatrisata, one finds scattered through the scalp irregularly shaped patches of alopecia whigh in the first stage may show perifollicular crythema but in the late stage show smooth atrophy of the skin without any signs of finlinimation. The loss of hair is

permanent

Histopathology. In the arly stage, one finds a predominantly perifollicular infiltrate composed almost entirely of lymphocytes and a few histocytes. The infiltrate is present around the upper and the middle thirds of the follicles and spares the lower third. It penetrates into the walls of the follicles and into the sebaceous glands, but not into the lumina of the follicles. Follicular hyperkeratosis may be present (Miescher and Lenggenhager). Gradually, the infiltrate de stroys the follicles and the sebaceous glands.

In a fate lesion, the epidermis is atrophic and the dermis shows fibrosis. The follicles and the sebiceous glands are absent, but arrec

tores pilorum and sweat glands pre, at least in part, preserved

Differential Diagnosis. Invocond lupus crythemitosus, the inflam matory infiltrate not only is located around hairs and sebaceous glands but also is distributed in a patch; fashion throughout the dermis In addition, the epidermis shows liquefaction degeneration of the basal cell layer, hiperferances; and more marked lenauric plugging not limited to the follicles. In the late cicarrical stage, a differentiation of the two diseases may be impossible.

Folliculius decalvans, which represents a folliculitis of the scalp, differs from alopecia cicatrisata by showing formation of intrafollicular abscesses (pustules). The perifollicular infiltrate often contains a fair number of phrsma cells which are absent in alopecia cicatrisata.

(Miescher and Lenggenhager)

PURPURA

Purpura represents a hemorrhage into the skin Lesions less than 3 mm in diameter are called petechiae Larger lesions are called ecchymoses

non which occurs in sensitized animals at the site of a subcutaneous injection of foreign protein. The presence of feulocytoclasis in ana phylacoid purpura would therefore support the ties that this disease represents an affergic reaction (Ruiter and Brandsma).

PURPURA PIGMENTOS \ PROGRESSIVA (Majocchi Schamberg)

Four diseases are included under this term purpura minilaris telangiectodes (Vajocchi) progressive piginentary dermatosis (Scham



Schamberg) \ group of capillaries located in a papilla show swelling prol leration and degeneration of their endothelial cells. \ \text{small} blocus of extraoastion of eighthcities can be seen to the right of the group of capillaries (x400)

herg) pigmented purpuric lichenoid dermatitis (Gougerot and Blum) and angioma serpiginosum. They are so closely related to one an other that often they cannot be differentiated on clinical or histologic present predominantly, but not exclusively around the damaged vessels (Fig. 60). It consists largely of neutrophils and varying amounts of cosmophils with only a few lymphocytes. A characteristic feature is the presence of many scattered nuclear fragments the result of the disintegration of neutrophils (leukocytoclasis). Fatravasation of

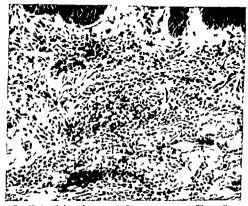


Fig. 60 Anaphylactoid purpura (inflammator) purpura). The capillaries show swelling and degeneration of endothelial cells. Viather severe inflammatory inflirate is present especially around the capillaries. It is composed largely of neutrophilis many of v high show disintegration of their nuclei (leukocytochisty.) This being an early lesion no extra is tion of erythrocytes has as yet occurred (×200).

erythrocytes is usually present although it may be absent in the early stage of the disease

In older lesions extravasation of red cells usually is a prominent feature. In addition, hemosiderin may be present due to the decomposition of red cells. The endothelial lining of the vessels may show proliferation. The infiltrate contains fewer neutrophils and consists predominantly of lymphocytes. However, nuclear fragments are still present, as a rule.

It may be pointed out that fragmentation of the nuclei of neutro phils is prominent in the histologic picture of the Arthus phenome non which occurs in sensitized animals at the site of a subcutaneous injection of foreign protein. The presence of leukocytoclasis in ana phylactoid purpura would therefore support the view that this disease represents in illergic reaction (Ruiter and Brandsmi).

PURPURA PIGMENTOSA PROGRESSIVA (Majocchi Schamberg)

Four diseases are included under this term purpura annularis telangiectodes (Majocchi) progressive pigmentary dermatosis (Scham



Fix 61 Putjura jugmentosa progressiva (Majocchi Schamberg) \ group of capillaries located in a papilla show suelling prodiferation and degeneration of their endothelial cells \ small focus of cruza-austron of cryption cytes can be seen to the right of the group of capillaries (x400)

berg) pigmented purpure lichenoid dermatitis (Gougerot and Bluin) and angioma serpiginosum. They are so closely related to one an other that often they cannot be differentiated on clinical or histologic

grounds Their separation into different entities, therefore, is un warranted (Randail, Kierland and Montgomery) The term purpura pigmentosa progressiva is suggested for this disease

The basic process is a chronic capillaritis of unknown cause occur ring in the upper dermis and leading to capillary fragility. Clinically, the primary lesion consists of purpuric puncta appearing in groups and slowly extending so that various sized patches form. Gradually, pigmentation, due to the deposition of hemosiderin, supervienes and in cases of long standing, may dominate the clinical picture. Inflummatory signs (such as crythema, scaling and papules) may be present or absent. In most instances, the disease is limited to the lower extremities but may be more or less generalized. The disease, though chronic, is harmless.

Histopathology. In early lesions, the capillaries of the upper dermis show swelling degeneration and proliferation of their endothelial cells. Often the number of capillaries appears to be increased. Small amounts of extravasited red cells are usually found in the vicinity of some capillaries. A cellular infiltrate, consisting largely of lymphocytes together with some histocytes and occisionally a few neutro phils is present in the upper dermis, especially in the vicinity of the capillaries. The inflammatory infiltrate may invade the lower epidermis and provoke mild liquefaction degeneration of the basal cell layer, mild spongiosis of the stratum malpighii and patichy para keratosis.

In older lesions, the number of capillaries is usually increased Some show proliferation of their endothelium and others dilutation of their lumen Extravasated red cells may no longer be present, but hemosiderin is almost always found, though in varying amounts. The infimmatory infiltrate is less pronounced than in the early stage. The epidermis may show slight atrophy with absence of the reteriodges.

Differential Diagnosis It may be difficult to differentiate purpura pigmentosa progressiva histologically from stasis derimation because inflammation extravasation of erythrocytes and deposits of hemo sideriu occur in both However the process extends much deeper in stasis derimatitis and in addition fibrosis of the derims and fibrous thickening of the walls of medium sized vessels in the lower derims is commonly present (see page 73). Anaphylactoid purpura differs from purpura pigmentosa progressiva by the predominance of neutrophils in the infiltrate and the presence of leukocytoclasis (see page 128).

RELAPSING FEBRILE NODULAR NONSUPPURATIVE PANNICULITIS (WEBER CHRISTIAN DISEASE)

This disease is characterized by the appearance of crops of indurated, tender nodules and plaques in the subcutaneous fat. As the lesions involute, they often leave a depression in the skin. The overconductions as a rule, no involvement other than mild crythema.

Histopathology. Histologically, the disease can be distanced three stages. The first two stages occur while there is inducation clinically During the third stage, depression of the skin develops. The first stage is observed only rarely because it is of short duration it is probable that it does not always occur. Most sections show a combination of the changes of the second and the third stage.

In the first stage (acute inflammatory stage), there is, between the fat cells an inflammatory infiltrate composed of polymorphonuclear leukocytes, lymphocytes and firstocytes Polymorphonuclear leuko

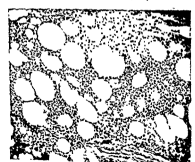


Fig. 62 Relapsing februle nodular nonsuppurative tor

cytes may predominate (Fig. 62) (Ungar Lever) Abscess formation does not occur

In the second stage (macrophagic stage) the infiltrate consists pre dominantly of histocytes A few lymphocytes and plasma cells are present. The histocytes are seen inviding and digesting the fat cells Such histocytes (macrophages) are large and have a formy cytoplasm



Fig 63 Relapsing febrile nodular nonsuppurative panniculitis Second and third stages. The left side and the center of the field show the second stage foam cells (macrophages) invading and digesting the fat cells. The right side shows the third stage re plicement by fibrotic connective tissue (×100)

(Fig. 63) Some of them are multinucleated. In some areas numerous macrophages with formy cytoplasm completely replace the fat cells

In the third stage (fibroblastic stage) fibroblasts intermingled with lymphocytes replace the macrophages. Collagen is laid down with resulting fibrosis

The epidermis and the dermis show no involvement. In some cases the subcutaneous vessels show pathologic changes such as edema and thickening of their walls (Cummins and Lever Tilden Gotshalk and Avakian)

In cases of liquelying panniculitis (Shaffer Binkley) the second macrophagic stage instead of being followed by fibrosis is followed by Inquefaction of the foam cell infiltrate. One finds an amorphous foams matrix in which the nuclei of foam cells as well as lymphocytes and some polymorphonuclear leukocytes are suspended.

It is now established that the disease may affect the internal adipose tissue. Four autopsies are on record. In three of them. internal lesions of pamiculitis were present which had the same bistologic appear ance as the cutaneous lesions. Involvement was slight in one of these cases (Spain and Foley) but extensive in two (Ungar Mostofi and Engleman). However only one of these patients (Mostofi and Engleman) However only one of these patients (Mostofi and Engleman) are sufficiently as a result of this disease.

Differential Diagnosis The histologic uppearance of nodular pan inculits is diagnostic in the second stage because there is no other condition in which one finds such preponderance of form rells in the sali cutineous fat. In foreign body granuloma one usually finds be sides form cells a great variety of other cells including foreign body structells.

SCI ERFMA NEONATORUM

Two types occur generalized sclerema neonatorum and nodular sclerema neonatorum (subcutaneous fut necrosis of the newborn adiponecrosis subcutanea neonatorum lipopliagic granuloma)

In generalized sel rema neonatorium, the skin of the entire body bis a navlike apperiance and is hard, dry and cold. Death usually actus within a teel. In nodular selerana neonatorium one observes deep seated industated areas in the subcutaneous fat. The process is localized and self limited, the lessons requiring about 4 months to disappear. The general health is not affected.

Histopathology The fundamental lesion is the same in generalized scientian neonatorium and in nodular sclerenia neonatorium (Zeek and Madder Flory) and both may occur simultaneously (Eichenlaub and Sindler). One finds degeneration necrosis and crystallization of the subcutaneous fai together with an inflammatory reaction and fibross. It is possible that the disease is due to a delay in the maturition of the fat and that on account of this there is an unusually low obecacted content of the fit so that it solidifies more easily than normally Sooth Pace and Paus).

GIVERALIZED SCHEFMA VIOLATORUM. The fat cells of the subcutis vari greath, in size and shape. In some areas they are necrous and have lost their outling. Needle-shaped empty, clefts he singly, or in ridiral arrangement inside of fat cells (Fig. 64). In frozen sections these clefts are it und to be occupied by crystals. The crystals fail to stain with fat stains Some of them are doubly refractite in the polarizing microscope (Reich. Zeek and Madden). The chemical composi-

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tion of the crystals is not yet established fully. Most investigators regard the crystals as those of fatty acids (McIntosh). A granulomatous infiltrate is present mainly at the periphery of the fat lobules but also in scattered foci throughout the lobules. The infiltrate is composed of lymphocytes, histocytes, foreign-body giant cells and fibroblasts. The septa between the fat lobules are considerably thickened.

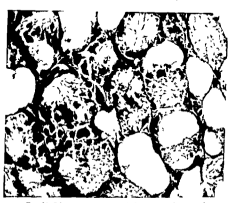


Fig. 64 Sclerema neonatorum (subcutaneous fat necrosis of the newborn). Severil fat cells contain needle shaped clefts in a radial arringement. These clefts are indicative of fit crystals. The fat crystals themselves are not visible because of fivation in Helly's solution, which dissolves fat. A granuloma tous infiltrate continuing foreign body, giant cells is present between the fat cells. (x 100)

In two cases of generalized sclerema neonatorum, autops, revealed in the visceral fat lesions which were histologically identical with those of the subcutaneous fat. In one case (Zeek and Madden), the lesions were widely distributed. In the other case (Flory), they were limited to the perirenal fat and to the fat about the ribs.

NODULAR SCIERTMA NEONATORUM (SUBCUTANEOUS FAT NECROSIS OF THE NEWBORN). One observes, as a rule, a more marked inflammatory reaction and a much larger number of foreign body giant cells than in generalized sclerema neonatorum (Fig 64) The foreign-

body giant cells may contain fat crystals (Fox). Healing takes place with fibrous

Differential Diagnosis The Instologic picture of sclerema neona torum differs from that of relapsing februle nodular nonsuppurative pannicultus by the presence of fat crystals and the absence of foam cells

HEREDITARY EDEMA OF THE LEGS (MILROYS DISEASE)

This disease occurs as a solid white indolent and persistent edema of the lower extremities. Usually several members of a family are affected.

Histopathology The dermis and especially the subcutineous fat show severe interstitual edemi. There is in increase in the amount of collagen in the dermis and segmentation of the subcutineous fat by thick strands of collagen. The capillaries and the lymphatics are dilated and increased in number and show slight perivascular lym lhocyte infiltration. The walls of the larger blood vessels may be thickened by fibrosis.

CHONDRODERMATITIS NODULARIS CHRONICA HELICIS

In this disorder one or several small well-defined hard painful indules are found on the upper mitigin of the ear. The surface of the modules often is hyperkeratotic. After removal of the keratotic layer a small lifter may be systulized.

Histopathology The epidermis shows hyperkeratosis irregular translussis and usually also central ulceration. The dermis and frequently also the perichondrium are permeated by a chronic inflammators, gradulomatous infiltrate composed of lymphocytes, plasma, and the composed of lymphocytes.

and secents. There is called the carriage instead of statung blue stains homogeneously pink with hematoxylin and enan in the affected areas.

The pathogenesis is not clear. Whereas most authors regard dekernine changes in the cattilage as the primary change (Foerster Fhemis). Newcomer recently has pointed out that similar degenerative changes occur in the aural cartilage with advancing age. It is probable that traium such as frostbate or pressure causes focal degeneration in the details of the ear. Due to poor sacularization

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in the affected area, repair does not ensue and the lesion becomes a focus of chronic inflammation.

SCABIES

Scabies, which is caused by the itch mite. Acarus scabies, presents burrows as its characteristic lesion. The burrows, produced by the female mite, occur on the palmar surfaces of the hands and the lingers,

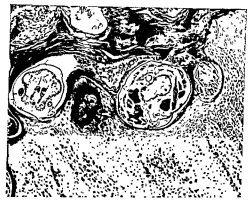


Fig. 65 Scabies (Norwegian scabies) Two female inites are located entirely within the horny layer of the epidermis (×200) (Robert N. Buchanan, Jr., M.D.)

on the interdigital skin, on the flevor surfaces of the urists, on the unpiles of women and on the genitals of men. They appear as fine, angular or tortuous black, is threads, a few millimeters long. Often a vesicle is visible near the blind end of the burrow. In addition to the burrows, scabies presents a papular eruption which is usually most pronounced on the abdomen, the lower portions of the buttocks and the anterior axillary folds.

In especially susceptible individuals, the clinical picture of so called sorwegian scabies may result, showing diffuse hyperkeratosis, scaling and crusting of the skin and burrows everywhere on the skin, even on the face and the scalp

Histopathology Histologic examination reveals the burrow in almost the whole of its course languard to the lucroy laser. Only the extreme blind end of the burrow is either in contract with or extending the textum malpighi. The femile mite is siturted it the blind end of the burrow (Fig. 6.) Its head is bored more or less deeply into the statum malpighia. Intracellular and interellular edema is present in the stritum malpighia beneath the mouth parts of the mite to such extent that vesicle formation often results. The mite thus takes its food in fluid form. The deems beneath the burrow shows a strong inflammatory infiltrate which is composed predominantly offinmphorytes.

Norweginn scables shows a much greater number of burlows than ordinary scables so that almost every section shows several of them (Ingram)

INSECT BITTS

Insect bites especially when caused by ticks may cause persistent lesions which pose a diagnostic problem clinically as well as histo-logitally. The clinical appearance of the lesions usually is that of a firm papile of of a nodule with or without central ulceration.

Histopathology. The epidermis frequently shows pseudo epitheliomitous hyperplasia (see page 331). The dermis presents a dense granulomatious infiltrate which often extends into the subcutaneous tat. It consists principally of cosmophilis and plasma cells admixed with himphoxytes and instructures some of the histocytes occasionally shown under that are hyperchromatic show mutout figures or are by nucleated. In some lexions large lymphoid follocles with germinal criters are observed (Winer and Strakosch Allen). In rare instances parts of the insect are found in the dermis either free in the tissue automided by a foreign body reaction or within an epidermal in clusion cast (Allen).

Differential Diagnosis. If parts of the insect are present in the section the diagnosis is obvious. In their absence the true nature of the lenon may be missed earth. The pseudo epitheliomatious hyperplasis must be differentiated from squirmous-cell carcinoma (see page 1.2%). The present of minote figures in the historytes together with the large, number of cosmophils may suggest incosis fungiodes the presence of lamphoid folibiles may suggest hyposis fungiodes the presence of kimphoid folibiles may suggest hyposis fungiodes the brune leared historytes if present require differentiation from the Sternberg Reed cells of Hodgkins discase However the abundance of pluma cells in susception with the eosimophuls usually rules out lymphoma.

LICHEN URTICATUS (PAPULAR URTICARIA)

This is a recurrent, pruritic eruption occurring, especially in children, during the summer months. The lesions consist of edematou

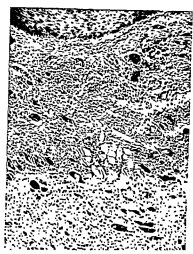


Fig. 66 Foreign body granuloma caused by a silk suture. The silk suture is located in the center of the field. Around in there is a severe influmentory infiltrate containing numerous foreign body grant cells (×200).

papules Sensitivity to flea and bedbug bites is now generally regarded as the usual cause (Goldman; Shaffer, Spencer and Blank).

Histopathology. The appearance is nonspecific Acanthosis and, in some cases, hyperkeratosis are present. The stratum malpighi shows intercellular as well as intracellular edema. A rather pronounced chronic inflammatory infiltrate is present around the vessels of the upper dermis.

FOREIGN BODY GRANULOMAS

Many foreign substances when injected or accidentally implanted into the skin may cause a foreign body reaction. In addition, certain



Fig. 67 Lipid granuloma caused by mineral oil (parafinoma). The many large and small ovoid or round cavities which give the section a Suiti cheese, appearance represent spaces filled with mineral of (parafi n) (x200).

substances formed within the body may produce a foreign body reaction when deposited in the detriits or in the subcutaneous are a Fxamples are the f

epithelioma o

(33)5

Histopathology \(\) typical foreign body reaction shows around the foreign material histocytes and foreign body giant cells (for

their description, see page 36) In addition, lymphocytes and plasmicells are present (Fig 66) The histocytes and the giant cells often exhibit phagocytic activity and then contain some of the foreign material

Some foreign body granulomas have a rather specific appearance Among them are lipid, tattoo, silicon and beryllium granulomas

LIPID GRANULOMA (PARATFINOMA)

Granulomas following injections of oily substances such as mineral oil (paraffin), cotton oil, or camphor oil occur as irregular, hard, nodular substances such as a literature para des despensables.

nodular subcutaneous swellings. Ulcoration may develop
Histopathology. Histologically, virind granuloms have a Swiss
cheese' appearance because of the presence of numerous ovoid or
round cavitics (Fig. 67). These cavities represent spaces occupied by
the oily substance (Conrad and Weiss). The spaces between the cavities are taken up by an infiltrate composed largely of dymphocytes
and some plasmi cells. In addition, there are groups of histocytes
some of which, due to the ingestion of lipid, have a formy cyto
plasm. Variable numbers of foreign body grant cells are present. In
Adder lesions-afforosis is prominent

Under the term sclerosing lipogranuloma, Smetani and Beinhard have reported as a post traumatic process subcutaneous granuloms with the same histologic picture as just described. However, Best Mason, DeWeerd and Dahlin have thrown serious doubt on the existence of such an entity. Investigating two similar cases, they found by chemical analysis that the lipid material was not body fat but mineral oil. The lipid material in their cases stained with Sudan 4 but not with osmic acid.

TATTOO GRANULOMA

Histopathology. Ordurary tattoos show diffusely scattered granules in the upper half of the dermis within phagocytes and lying free in the tissue, without any inflammatory reaction

However, if an inflammatory reaction occurs, due to allergy, the inflammatory infiltrate shows aside from the phragocytes numerous lymphocytes with an admixture of many, cosinophils and a few plasma cells (Rostenberg Brown and Cruo)

SILICON GRANULOMA

Silicon may be introduced into the skin through the contamination of lacerations with particles of soil. Such wounds heal at first and then, many months or years later indurated nodules develop in the skin or the subcuttneous tissue.

Histopathology. The histologic picture is indistinguishable from that of sarcoidosis (see pige 188) except for the presence in some of the grant left of colorless, spiculated, crystalline particles varying in the process of trade trible to 100 microns in length. When examined

BERYLLIUM GRANULOMA

Beryllium gruntloms of the skin may form in two different ways (Grier Nash and Freiman). They may arise as a manifestation of



to 6 Berjilium granuloma caused by Inceration with a fluorescent light tube. There is a large stee of spectrum necross surrounded by tubeculoud granulation issue. In the other several surrounded by tuberound granulation issue.

viscenic hersiliosis, in which case it must be assumed that particles of hersilium recicled the skin through the circulation, or they may develop following adactration of the skin through which beryllium entered the tissue. Such lacerations were observed some years ago from cuts with fluorescent help half.

their description, see page 36) In addition, lymphocytes and plasma cells are present (Fig 66). The histocytes and the giant cells often exhibit phagocytic activity and then contain some of the foreign materials.

Some foreign body granulomas have a rather specific appearance Among them are lipid, tattoo, silicon and beryllium granuloms

LIPID GRANULOMA (PARAFFINOMA)

Granulomas following injections of oily substances such as mineral oil (paraffin), cotton oil, or camphor oil occur as irregular, hard nodular subcutaneous swellings. Ulcoration may develop

Histopathology. Histologically, vilpid granulomis have a Swiss cheese" appearance because of the presence of numerous ovoid or round cavities (Fig. 67). These cavities represent spaces occupied by the oily substance (Contrad and Weiss). The spaces between the cavities are taken up by an infiltrate composed largely of lymphocytes and some plasma cells. In addition, there are groups of instrocytes some of which, due to the ingestion of lipid, have a formy cyto plasm. Variable numbers of foreign body grant cells are present. In Adder lesions-fibrosis is prominent.

Under the term selerosing hipogranuloma, Smetana and Bernhard have reported as a post traumatic process subcutaneous granulomas with the same histologic picture as just described. However, Best, Mason, DeWeerd and Dahlin have thrown serious doubt on the existence of such an entity. Investigating two similar cases, they found by chemical analysis, that the 'lipid' material was not body far but mineral oil. The 'lipid' material in their cases stained with Sudan 4 but not with ownic acid.

TATTOO GRANULOMA

Histopathology. Ordinary tattoos show diffusely scattered grammles in the upper half of the dermis within phagocytes and lying free in the tissue, without any inflammatory reaction

However, if an infimumatory reaction occurs due to allergy, the inflammatory infiltrate shows aside from the phagocytes numerous lymphocytes with an admixture of many cosmophils and a few plasma cells (Rostenberg Brown and Caro)

SILICON GRANULOMA

Silicon may be introduced into the skin through the contamination of lacerations with particles of soil. Such wounds heal at first and then, many months or years later inducted nodules develop in the skin or the subcurneous tissue.



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Acrodermantis Communa

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nodules over which the skin remains intact. They are always few in number The most serious lesion in systemic berylliosis is the chronic progressive pneumonitis which is frequently fatal

The cutaneous granulomas following laceration show as first sign incomplete healing of the laceration, followed by swelling, induration and tenderness and, finally, central ulceration

Histopathology. The cutaneous granulomas of systemic berylhosis often are indistinguishable from sarcoidosis (Hardy and Tabershaw) Occasionally, however, a moderate amount of caseation necrosis is present in some of the epithelioid cell islands. The quantity of beryl hum present in the lesions is too small to be demonstrable by spectrographic analysis

The cutaneous granulomas following Vaceration show, as a rule, , more caseation necrosis than the granulomas of systemic berylliosis The caseation necrosis not only may be present within some of the epithelioid cell islands but also may affect the entire center of the lesion (Fig. 68) (Neave, Frank and Tolmach). A collar of lymphocytes often surrounds some of the epithelioid cell islands, giving them the appearance of true tubercles Occasionally, Schaumann bodies, just like those of sarcoidosis, are present (Grier, Nash and Freiman) The epidermis shows acanthosis and may show ulceration. In some cases the presence of beryllium has been demonstrated by spectrographic analysis (Dutra)

SWIMMING POOL GRANULOMA

Abrasions in swimming pools may lead to circumscribed areas of nodular infiltration on the face resembling lupus vulgaris (Heller strom) or to granulomatous, vertucous lesions on the knee or else where resembling tuberculosis verrucosa cutis (Rees and Bennett) The lesions usually heal within 3 to 9 months

Histopathology. The histologic picture is similar to that of lupus vulgaris (Hellerstrom) or of tuberculosis verrucosa cutis (Rees and Bennett) Hellerstrom regarded his cases as instances of inoculation lupus vulgaris but moculations of tissue into guinea pigs have given negative results in all reported cases and Hellerstrom found acid fast bacilli in the tissue in but one of his six cases. Thus a tuberculous cause is not established Silicon granuloma is also unlikely because no crystalline particles were ever observed. The cause remains ob scure

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[\ \[\ 80 1761 1923 Swimming Pool Granuloma

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Eruptions Due to Drugs

Allergic reactions to drugs may cause various eruptions identical in their clinical appearances to cutaneous diseases occurring also as idio pathic entities. Drugs may cause, for instance, urticiria, erythemi multiforme, erythema nodosium, dermatitis, including generilized exfoliative dermatitis, purpura, folliculitis and periarreritis nodosi. The histologic picture is the same in these diseases whether they are due to a drug or occur in their idiopathic form.

Only histologic changes more or less specific for eruptions due to

drugs will be discussed

FIXED DRUG ERUPTION

Fixed drug eruptions are circumscribed lesions which persistently recur at the same site at each administration of the allergenic drug. The most common type of fixed drug eruption consists of one or several slightly elevated, erythematous plaques which on healing leave pigmented areas. Fixed drug eruptions may occur after the ingestion of phenolphthalein and coal tar derivatives such as amino pyrine and acetophenetidin.

Histopathology. There is an increase in melanin in the basil cells and in the dendritic cells of the epidermis. Melanin is present in the upper dermis. It is found predominantly in histocytes (melano phores) but also free in the tissue. The superficul capillaries are sur rounded by a mild inflammatory infiltrate composed of lymplicytes.

and histocytes (Weiss and Kile Chargin and Leifer)

QUINACRINE HYDROCHLORIDE (ATABRINE) DERMATITIS

Quinterine hydrochloride (Atabrine) may cause a subrette or a chronic dermattis, lichen plinus like lesions and exfoliative dermittiss (Bereston). In some patients the eruption is followed by anhibitorism in the involved areas.

Histopathology The histologic picture is usually that of a non specific chronic dermatius. In lesions that clinically resemble lichen plants, the histopathologic picture often resembles that of lichen

planus too As a rule however, the bandlike infiltrate is not so dense as in lichen planus and may contain eosinophils which are not present in true lichen planus (Wilson) in many cases one observes object-craosis with keratotic plugging of pilosebaccous follicles Numerous melanophores may be present in the dermis (Alden and Frank)

In patients in whom anhidrosis develops in areas of quinacrine hydrochloride dermatitis one may observe attrophy of the dermal portions of sweat ducts and dilatation and attrophy of the sweat glands Focal inflammation may be present about the sweat glands (Sulbberger Hermann and Zak)

EXPOLIATIVE DERMATITIS DUE TO DRUGS

The most common drugs to cause exfoliative dermatitis are arsphenamine gold salts the sulfonamides and phenobarbital

Histopathology The exfoliative dermatitis caused by these drugs has the same histologic appearance as exfoliative dermatitis due to other causes (see page 74 and Fig. 27)

Changes in the internal organs may be present such as interstitual injocerdatis (Brown and McNimara French and Weller Winer and Baer) interstitual nephitatis (Winer and Baer) and fatty degeneration of the liver with inflammatory infiltration about the portal canals (Winer and Baer).

BROMODERMA

Prolonged ingestion of bromides may cause the formation of granulomatous vertucous plaques which are called bromoderma. They occur usually on the lower extremities

Histopathology There are prollomatosis and considerable down ward proliferation of the epidermis often of such degree is to produce the picture of pseudo-epithehomatous by periode and the processing of the proc

The dermis shows an extensive granulomatous inflittate which may reach down into the subcutaneous layer It is composed of a great variety of cells including lymphocytes plasma cells and histocytes reutrophils are usually numerous and abscesses may be found seat tered through the inflittate Eosinophils are few or absent. The blood vessels are increased in number are dilated and show proliferation of their endothelium. Small areas of hemorrhage are often seen within the inflitrate

Differential Diagnosis The histologic picture of bromoderma is suggestive but nor diagnostic Pyoderma gangrenosum may show at

the margin of an ulcer, an identical histologic picture, including intra epidermal abscesses. Intra epidermal abscesses are also observed in older lesions of pemphigus vegetans and in blistomycosis. Pemphigus vegetans differs from bromoderina by the large percentige of cosmophils in the intra epidermal abscesses and in the granulom



Fig. 69 Bromoderm: There is downward proliferation of the epidermis. A large intra-epidermal thicker is shown. The dermis contains a granulomatous infiltrate (×50).

tous infiltrate (see page 80). Blastomycosis is easily differentiated by its numerous grant cells and the presence of yeast cells in them

IODODERMA

Iododerina is characterized by granulomatous lesions which may have the same clinical appearance as bromoderina. As a rule, however, there is less verrucous proliferation and greater tendency to infernation.

Histopathology. Histologically iododerma differs from bromo derma by showing less epithelial proliferation. The granulomatous infiltrate frequently destroys the epidermis resulting in ulceration. The infiltrate may be composed predominantly of histocytes some of which may show mitotic figures and hyperchromatic nuclei, so that

distinction from lessons of mycosis fungoides in the tumor stage may be almost impossible (Filer and Fox, Hollander and Fetterman)

ARGYRIA

This condition caused by prolonged ingestion of silver salts or their local application to the mucous membranes, is characterized by



Fit 70 trgitts Silver granules are present in the membrana proprise of the swell stands in some places, the granules are so dense that they form a solid black band (x400)

blash $_{\alpha}r_{\alpha}r_{\beta}$, sine colored discoloration of the entire skin-most pronounced on the exposed portions of the skin

Histopathology Silver is found in the dermis predominantly extractilitates white small tound brownish particles of authorm size it is never seen in the epidermis or its uppendings. The silver partacks measure less than I micron in diameter and he singly as well is in clumps. Mitrough visible in rotuning strains they appear more clearly in sections stained bybols with polychrome methylene blue. However, the best method for the demonstration of silver granules as he did both dilumination. If sections we pitced under a dark field microscope the silver appears as brilliantly refractile, white granules against a dark background. Many more granules can be seen than with direct illumination.

The silver grunules are most numerous in the membranae propries surrounding the sweat glands (Fig. 70) and in the subepidermal dermis. In addition silver particles may be seen in the connective tissue sheaths about the hair follicles and the sebaceous glands in the wills of blood vessels particularly their intima and adventitia in the arrectores pilorum and the nerves and diffusely scuttered through the middle and the lower dermis. Elastic tissue stains reveals a predilection of the granules of silver for elastic fibers. The location of silver in elastic fibers explains the presence of fingerlike chains of granules projecting into the papillary bodies (Hill and Montgomery). In many cases one finds increased amounts of melanin in the basal layer of the epidermis and melanin laden chromatophores in the dermis.

Silver is deposited not only in the skin but also in internal organs. It is found particularly in the intima of blood vessels and in the connective tissue of the internal organs. Analogous to the marked involvement of the basement membrane of the swear glands the basement membranes around the acini of the testes and of the choroid plexus are particularly rich in granules (Harker and Hunter).

Differential Diagnosis Histologic differentiation of argyra from other kinds of pigmentation is made readily. Melanin and hemo siderin possess larger granules which he to a great extent intracellularly in chromatophores and are nonrefrictile with dark field illumination. In pigmentation due to mercury, whether from the use of creams or in tatoo marks, the mercury is deposited in large coarse granules, throughout the epiderium and the derinis without any special affinity for the membrana propriato of the sweat glands. For differentiation from chrysiasis see below.

CHRYSIASIS

In chrysiasis which may follow the parenteral use of gold salts the skin of the exposed parts shows an ash gray discoloration

Histopathology Gold granules are light refractile with dark field examination like silver granules but they are larger and more are fur in size than silver granules. In contrast to silver granules they are found predominantly within cells. They lie in the endothelial and the perithelial cells o capillaries and in macrophages through out the upper dermis. Only occasionally do granules he free in the tissue spaces (Schmidt). In some instances gold particles are found in the basal cells of the epidermis (Kochs).

ARSENICAL KERATOSIS AND CARCINOMA

Prolonged ingestion of inorganic arsenic frequently produces multiple arsenical Leriatoses which may progress into squamous-cell carentoma. Recently it has been suggested that not only multiple cutaneous cancers but also multiple internal cancers may be caused (Sommers and McManus). Occasionally, multiple superficial basal cell epitheliomas form in addition to arsenical keratoses (Anderson Monteomers, and Waisman).

Arsenical keratoses resemble senule keratoses in their chincal appearance. They may occur anywhere on the skin but are found most frequently on the palms and the soles in contrast with senule keritoses which predominate on the face and on the dorsa of the hands.

Histopathology In early arsenical keratoses one observes hyper keratosis associated with acanthosis and irregular downward pro-liferation of the rete tridges. There usually its some degree of disorder of the squamous cells and pyknosis of some of the nuclei. The histories incluse this top some of the tridges within the tridges within the tridges.

In more advanced lesions in addition to the above mentioned

of the ouncer man in lowers disease. Although vacuolization of cells occurs also in Bowens disease this feeture is not so prominent as in assential keratoris. The presence of numerous vacuolated cells may be regarded as diagnostic of aisenical keratoris (Mont gomets) and Waisman). The vacuolated cells are twice or three times as large as normal squamous cells and possess small irregular deeply st immig nuclei. They resemble the Paget cells of Paget's disease of it empthe except that intercellular priceles are tsurily present it empthe except that intercellular pricels are tsurily present. I timately through invasion of the dermis frank squamous cell.

Ultimately through invasion of the dermis frank squamous cell carcinoma may develop (see page 329). Even in the invading type of carcinoma vacuolization clumping and dysteratosis remain prominent.

RIRI IOCD ADUS Fixed Drug Eruption

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Degenerative Diseases

SINIT DEGENERATION

Sende changes in skin not regularly exposed to helic manifest themselves chinically in thinning of the sin and atrophy of the sin cutinious fat. In exposed skin the changes usually are much more pronounced by in in skin that is not exposed and include marked

skin after minor trauma eccletimoses may occur (semile purpura) xposed to light semile retejadees and atrophy changes in the collagen

or in the Classic ussue are absent. Vild obliterative changes may be found in some of the vessels (Hill and Montgomery)

In sking I used to suntishit two types of degenerative changes are seen which I owever represent one and the same process. Dasophila degeneration of the collegen and setule elastosis. These changes hunted to the upper third of the derius may start as early as in the third decide of life and progress with age.

In Prophile de eneration, the collagenous fibers appear broken up into amorphous clumps and grundes which stain is on a second of the collagenous fibers.

on this observes in the upper derims separated from a somewhat in split epiderims by a narrow band of normal collage masses of twisted that. Birds straining fibers (Fig. 71) Because these fibers give the same atmosp recitions as elastic tissue. Unna referred to them as collision thinking they were the result of a merging of collagen with elastic fibers. However recent studies by x-ry diffraction and electron microscop's suggest that the essential change in senile elastions is degeneration of collagen fibers. Furthermore the degenerated miteral can be remoted—in contrast with elastic fibers—by treatment of the sections with tripsin and at the same time the

RIRI IOCP IDIN

Fixed Drug Truption

Chargin L. and Leifer W. Fixed eruntions due to the arsphenamines. I Invest Dermat 9 449 1040

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xerotica obliterans formerly regarded as independent diseases prob ably are identical with lichen sclerosus et atrophicus (see below)

Histopathology In lichen sclerosus et atrophicus one observes (1) hyperkeratosis with keratotic plugging (2) atrophy of the stratum



keratoris with follicular plugging, acrophy of the stratum malp ghu marked simphedema of the upper dermis and an inflammatory infitrate in the mid dermis. The edema in the subepidermal dermis is so marked that a bulla has resulted. (X100)

The hyperkerators is so marked that often the horny layer is thicker than the attophic stratum malpinghin. The stratum malpinghin is reduced to a few layers of flattened cells. The cells of the beasal layer show hydropic degeneration. The reterindges often are completely

158 Degenerative Diseases

elastic sturning properties of the skin disappear (Tattersall and Seville) This suggests that the increase in ordein staining material is caused not by an increase in the amount of elastic tissue but by a false, staining reaction of degenerated collagen Staining of frozen sections with fat stains occasionally reveals numerous fine lipid drop lets in the areas of basophilic degeneration or senile elastosis (Weidman Percival Hannay and Duthie)



Fig. 71 Senile degeneration (senile elastosis). In the upper detries septented from the cytderm's by a narrow band of normal collagen there are masses of twisted thick degenerated collagen fibers staining black with orcur-just like elastic tissue (X100).

Differential Diagnosis For differentiation from pseudoxanthoma elasticum see page 56

LICHEN SCLEROSUS ET ATROPHICUS

This disorder is characterized by flat topped white papules which coalesce to form white patches without any infiltration. The surface of the lesions often shows comedo like plugs. Occasionally, the patches become bullous due to accumulation of fluid beneath the epidermis. Lichen sclerosus et atrophicus occurs not only on the skin but also on the vulva and the glans penis. Araurosis vulvae and balanus

super out but they may persist in some areas and even show some irregular downward proliferation. In such proliferations, hydropic

degeneration of the basal cells usually is pronounced Beneath the epidermis there is a broad zone of pronounced edema

Within this zone the collagenous fibers are swollen and homogeneous and contain only a few nuclei. They stain poorly with eosin and other connective tissue stains. This change does not represent sclerosis (as the name of the disease would imply) but lymphedema. The hydropic degeneration of the basal cells together with the edema of the sub epidermal collagen may lead to the formation of subepidermal bullac (Cottschalk and Cooper) These bullae may thus be classified as being due to degeneration of the basal tells (see Classification of Bullie page 66) The elistic fibers are sparse or even absent in the zone of edema (Nomland)

In the wild-dermis beneath the area of edema there is an infiltrate which usually is perivascular but at times assumes a bandlike forma tion It is composed almost exclusively of lymphocates. In lesions of long duration the infiltrate may have almost disappeared

KRAUROSIS VIII.VAE

\ satisfactors classification of the atrophic lesions of the vulva has not yet evolved and to a certain degree the vagueness of the term kraurosis vulvae is responsible for this. Although some authors still regard it as an entity (Wallace and Whimster) others regard it as synonymous with lichen sclerosus et atrophicus (Laymon)

The simplest classification of the atrophic lesions of the vulva that can be offered at present is

- 1 Senite or Presente Atrophy There is atrophy of the vulvar mucosa but nXstenosis of the vaginal orifice. There may be itching and due to scratching as ulvitis may result
- 2 TICHEN SCLEROSUS FT ATROPHICUS (KRAUROSIS VULLAE) CON siderable attophy with stenosis of the vaginal orifice is present. The lesions are whitish in color sharply demarcated and may extend to the inguinal folds and the peri anal region. Itching may be present Because the lesions have a whitish color on casual inspection they resemble leukoplakia but they lick the indutation observed in leuko plakes. There has been uncertainty as to whether lichen sclerosus et

fic * 6 Krautosis vuls ae flithen seletosus et atrophicus). There are edema of the upper dermis and a landlike inflaminatory infiltrate beneath it. In all tunn there is irregular downward problems on of the rete ridges with l'ulropic degeneration of the basal cells. The latter feature is typical of I then seletonus et attophicus and rules out leukoplakia (×100)

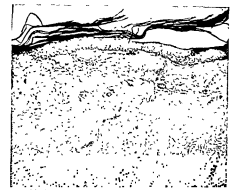


Fig. 73 Kraurosis vulvae (lichen sclerosus et atrophicus). There are hyperkeritosis atrophy of the stritum malpighii and mirked lymph edemi of the upper dermis with homogenization of the colligen (x100).



Fig 74. See facing page for legend.

upped out but they may persist in some areas and even show some irregular downward proliferation. In such proliferations, hydropic degeneration of the bisal cells usually is pronounced.

Beneath the epidermis there is a broad zone of pronounced edemic Multin this zone the collagenous fibers are swollen and homogeneous and contain only a few nuclei They stain poorly with coun and other connectine tissue stains. This change does not represent selerous (as the name of the disease would imply) but lymphedemia. The hydropic degeneration of the basal cells together with the edema of the subepidermal collagen may lead to the formation of subepidermal bullace (Cottschall, and Cooper). These bullen may thus be classified as being due to degeneration of the basal cells (see Classification of Bullace page 66). The elastic fibers are sparse or even absent in the zone of edepan (Nomland).

In the wind dermis beneath the area of edema there is an infiltrate which insually is peri actual but at times assumes a bandlike formation. It is composed almost exclusively of <u>hamblogues</u>. In lessons of long duration the infiltrate may have almost disappeared.

KRAUROSIS VULVAE

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The simplest classification of the atrophic lesions of the vulva that can be offered at present is

- 1 SENILE OR PRESENTE ATROPHY There is atrophy of the vulvar mucosa but mystenosis of the vaginal orifice. There may be itching and due to stratching, as ulvitis may result.
- 2 LICHEN SCLEROUS ET ATROPHICLS (REAUROSIS VULVAE) Con siderable atrophy with stenoses of the tagand ortice is present. The iessions are whiths in color sharply demarcated and may extend to the inguind folds and the peri and region leching may be present Because the lesions have a whitish color on casual inspection the resemble to the stage of the stag

Fig. 74 Arturents whise (lither selectories et atrophicus). There are edema of the upper dermis and a landlike inflammatory inflitrate beneath it hadd ton there is irregular down and productation of the reter ridges with hadropic degeneration of the 20-21 cells. The latter feature is typical of liches selectorie et arrophicus and rules out lexicophika (2/100).

atrophicus, a perfectly harmless lesion elsewhere, may progress into leukoplakia and squamous cell carcinoma when occurring on the vulva. Such instances recently have been reported (Wallace and Whimster), but it seems to be rare. Nevertheless, as Wallace and Whimster state, any state of atrophy occurring on the vulva has malignant potentialities.

3. LEUKOPLAKIA One or several whitish indurated plaques are present Leukoplakia may occur independently or secondary to senile atrophy of the vulva. Since leukoplakia is an early malignant lesion ("squamous cell carcinoma, grade ½," see page 328), development of squamous cell carcinoma is common.

Histopathology. The histologic picture of centle or presentle atrophy is nonspecific, consisting of atrophy of the epidermis and a mild to moderately severe chronic inflammators infiltrate in the

upper dermis

Vichen sclerosus et atrophicus of the vulva (kraurosis vulvae) shows the same histologic picture as on the skin with the exception that lesions located on the mucous membranes show nox cratotic plugs (Fig. 73). Irregular downward proliferation of the rete ridges in which the basal cells show considerable hydropic degeneration occur more often than on the skin (Fig. 74). They should not be confused with the early atypical proliferations of leukoplikia which do not contain such hydropic basal cells.

Deukoplakia shows atypicality of the epidermis with irregular downward proliferation of atypical cells (see page 328). The hydropic degeneration of the basal cells seen in lichen sclerosus et atrophicus

ıs absent

BALANITIS XEROTICA OBLITERANS

Balantis verouca obliterans represents lichen sclerosus et atrophicus of the glans and prepuce (Laymon). It is a chronic, progressive atrophic process which frequently eventuates in urethrabstenosis. In very rare instances carcinoma may supervene (Grutz)

Histopathology The histologic picture is that of lichen sclerosus et atrophicus Because of the absence of follicles in the areas of

involvement, no keratotic plugging occurs

STRIAE DISTENSAE

Striae distensae occur in pregnancy and obesity, and particularly in Cushing's disease. They represent linear areas of cutaneous acrophy Histopathology. In the early lesions, the clastic fibers are trans

Histopathology. In the carly lessons, the clastic fibers are trans formed into numerous faintly staining fibrillae due to fraying of the clastic fibers. In addition, there are a mild perivascular infiltrate and

a distortion of the collagen bundles 'Old lesions show almost complete absence of elastic fibers in the center. At the mirgin coiled and clumped elastic fibers are interspersed with fine, poorly staining fibriliae.

On the basis of these findings. Ebert has concluded that strize distensee are not due to mechanical tearing of elastic fibers alone but that degeneration of the elastic fibers precedes their disappearance in the center of the strize.

MACULAR ATROPHY (ANETODERMA)

Macular atrophy of Schweninger and Buzzi or anetoderma is characterized by atrophic oxal patches located particularly on the truni. The shin of the patches is thin and bluins white and bulges slightly. The lesions may give to the palpating finger the same sensa tion as does a hermal orifice Early lesions may show mild erythema. In the literature a difference has been made between a primary

In the interactive a difference has been made between a primary idiopathic type of macular attrophy and a secondary type developing in patients with diseases such as syphilis lupus crythematosus and acrodermatitis chronica atrophicans but in sites not clinically affected by these diseases (Scull and Nomland). It is likely that such association is merely coincidence.

11

les "a non at a many disappearance of the elastic tissue throughout the dermis. The collagen remains unaffected. Thus a diagnosis of this disease can be made only from sections stained for elastic tissue.

ATROPHODERMA RETICULATUM (FOLLICULITIS ULERITHEMATOSA RETICULATA)

The eruption which is limited to the sides of the face consists of numerous small closely set areas of atrophy separated by narrow ridges of normal skin resulting in a reticulated appearance

Histopathology The epidermis shows diminution of the number of reer ridges and follicular plugging. Horn cysts caused by keratum ration of hair follicles are present in the dermis. Sebaceous glands are few in number and small in size. In addition one observes degeneration of the collagen and fragmentation of the elastic fibers.

COLLOID MII IUM (COLLOID DEGENERATION OF THE SKIN)

Colloid milium is characterized by pinhead sized round sharply circumscribed yellowish nodules of the skin. The nodules have a

translucent appearance and on puncture give exit to a soft gehtt nous mass. The forehead is the site of predilection

In occasional instances instead of small nodules large plaquelike lesions are present. To such cases the term colloid degeneration of the skin lars been applied (Reuter and Becker).

Histopythology Histologically colloid milium is characterized by the presence of circumscribed masses of colloid material in the sub-

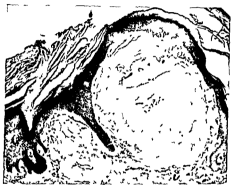


Fig. 75. Colloid milium. In the upper dermis sep rated from the epiderms by a narrow zone of normal collagen there are is a large round spaces incompletely file t in the fisture! I may of 1 magenes is material. This material represents colloid. The cold id contains a militarity material represents colloid.

epidermal region. It is assumed by many that colloid represents a form of collagen degeneration. Prakken has observed that collagen before becoming colloid passes through the stage of pasophilic degeneration. However, it is possible that colloid like amyloid represents a deposit rather than a product of collagen degeneration (Percival, and Duthie).

The epidermis shows hypetheritosis and atrophy of the stratum malpighit Large nearly ound spaces he close to the epidermis separated from it by only a narrow zone of normal collagen. These spaces extend through the upper third of the dermis and are sharply demarcated by collagen bundles arranged circularly around them.

They are filled incompletely by a fissured mass of homogeneous appearing material containing a moderate number of nuclei (Fig 75) The material referred to as colloid usually stains eosinophilic with hematoxylin and cosin though to a lesser degree than normal collagen Occasionally however it stains faintly bisophilic The fis. suring of the colloid is due to fixation and dehydration. The nuclei within the colloid are well preserved and represent fibroblasts Flastic tissue stains reveal clustic fibers within the masses of colloid however they are fragmented and fewer in number than in normal collagen.

Inspolloid degeneration of the skin colloid is present not in super ficial circumscribed foci bin diffusely throughout the dermis (Reuter

and Becket)

Differential Diagnosis Differentiation of colloid milium from handolosis (see page 274) requires special staining since these two substances greatly resemble each other morphologically and usually stain alike a pale pink with hematoxylin and eosin. One may use van Gieson's stain which stains colloid yellow and amyloid pink or methyl violet with which only amyloid is stained

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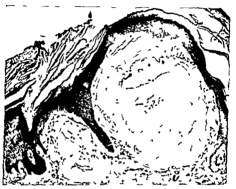


Fig. 75 Colloid milium. In the upper dermis separated from the epiderms by a narror zone of normal coll for there are two large round up ces incompletely filled i the r i sured mass of hamogeneous material. This material represents of lloid. The colloid contains a malerate number of nuclei (x/100).

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Bacterial Diseases

IMPETIGO
Impetigo may be caused by streptococci or by staphylococci Bullae are the primary lesion in either case. In the streptococcal variety, the bullae soon rupture leaving sharply demarcated erosions which be



with neutrophils is present. The epidermis shows spongiosis Many neutrophils are seen migrating through the epidermis (×200)

come covered with heavy, honey colored crusts. In the Yaphylococcal variety the builde are more durable and dominate the clinical pic ture Staphylococcal impetigo occurs especially in the newborn (for merly also called pemphigus neonatorum)

Histopathology In both the streptococcal and the staphylococcal variety of impetigo the bulla arises directly beneath the horny layer

166 Degenerative Diseases

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IMPETICO BOCKHART presents a subcorneal pustule situated in the opening of a hair follicle. The upper portion of the hair follicle is surrounded by a considerable inflammatory infiltrate containing second outpets of polymorphomaticiar leukocytes.

varying numbers of polymorphonicleir leukocytes
A furuncity presents on histologic extinuation a perifollitular
abscess composed of a deme mass of polymorphonicleir leukocytes
with a few lymphocytes. The hair with its follicle and its sebaceous

glands is destroyed in the process

In FOLLOWITS BARBAF the perifoliculus is as a rule less acute than in humanculous and does not necessarily result in a perifolilicular alvess Frequently the infiltrate around the follicle contrins relatively few polymorphomuclers leukocytes but consists mainly of hymphocyte plasma cells and histocytes Some of the sebaceous glands undergo destruction but the hair follicles may escape destruction Foreign body grun cells frequently are present around the hair follicles and the remnints of sebucous glands. In many cases instead of being limited to the vicinity of the follicles the infiltrate extends through the entire upper dermis.

FOLLICULITIS KEI OID ALIS (DERMATITIS PAPILLARIS CAPILLITII)

Folloculitis keloidalis represents a chronic folloculitis resulting in keloidal scarring It occurs on the nape of the neck, in men. In early cases one observes iodicular popules pustules and occasionally abscesses. The lesions are replaced gradually by fibrois nodules.

Anscesses The festions are replaced gradualty by librous nodules
Histopathology Early lesions show the same histopathologic picture as that of a furuncle Older lesions show chronic granulation

tissue concumnant yes and 6b around

ter in a comment the histologic pic ture is the same as in keloid showing bundles of hypertrophic and sclerotic collagen

HIDR ADENITIS SUPPUR ATILA

Hidradenins suppuration represents a chronic staphylococcal in territor of the apoctine glands in the avillary or the public regions Early lesions consist of red tender nodules, which become fluctuating and finally discharge pus. Ulcers may develop and healing takes place with considerable scarting.

Histopythology The infection enters by way of the follieles and excretory ducts of the apoctine glands. The earliest cellular reaction is encountered in the subcutaneous tissue within and around the

941

(see "Classification of Bullae," page 64) The bull contains fibrin polymorphonuclear leukocytes and some lymphocytes (Fig 76) With Gemsa's or Gram's stain, groups of coccl. can be recognized within the bulla. They may be extracellularly or within neutrophils

The stratum malpighi underlying the bulla shows spongiosis nyiny neutrophils can be seen inigrating through it The upper dermis contains a moderately severe inflammatory infiltrate of neu

trophils and lymphocytes

At a later stage, when the bulla has ruptured, the horny layer is absent, and a crust composed of fibrin and neutrophils may be found resting on the stratum malpighii

PRYSIPFI AS

Frysipelas is a localized acute inflammation of the skin caused by streptococci. It is characterized by the presence of well demarcated dusky red areas with or without edema and vesiculation. A typical sign is the advancing acd, raised and industed border.

Histopathology. The dermis shows marked edemy and dilutation of lymphatics and capillaties. When the edemy is intense subept dermal bullae may be present. There is a marked diffuse infilitate extending throughout the dermis and, occasionally, into the sub-cutaneous fat. It is composed cluefly of polymorphonuclear leuko-cytes and some lymphocytes. If sections are stained with Giemass or Gram's stain, streptococci are found chiefly in the lymphatics, but also distributed in the tissue.

FOLLICULITIS (IMPETIGO BOCKHART, 1 URUNCI F FOLLICULITIS BARBAE)

Pustular follicultus may occur in three forms as impetigo Bock hart, as furuncle and as follicultus barbae (sycosis vulgaris). All three are caused by staphylococci

are caused by staphylococci
Impetigo Bockhart represents a superficial pustular follicultus and
is characterized by an eruption of pustules many of which are pierced
by a hair

A furuncle is a deep seated folloulitis leading to a perifollicular

cellulitis and terminating in suppuration and necrosis

Folliculitis barbie is a deep serted folliculitis peculiar to the bearded region. At first there are follicular pipules and pustules which are followed by diffuse crythema crusting and infiltration of the skin. Abscesses may be present or absent.

Histopathology. These three discuses cannot always be differen

tiated on a histologic basis

Osler nodes are erythematous slightly raised tender intracuta neous nodes averaging 5 mm in size. They occur most frequently on the fingertips and as a rule last 4 or 5 days.

Janeway lesions are small macular or papular lesions measuring from 1 to 4 mm in diameter. They are usually red but may be par-tially hemorrhagic. They occur most commonly on the palms and the

soles Unlike Osler nodes they are never tender

Mistopathology The petechiae show the histologic picture of an inflammatory purpura (see prige 127) presenting severe vasculitis in addition to the extravastion of red cells. One observes marked endo thelial proliferation of capillaries leading to narrowing or even obliteration of the lumina An infiltrate composed of polymorphonuclear leukocytes lymphocytes and histocytes surrounds and in vades the capillaries Streptococci have never been found in the lesions (Merklen and Wolf Cornil Mosinger and Jouve)

The Osler nodes show involvement not only of the capillaries but also of the dermal arterioles and venules which show intense endo thelial proliferation and not infrequently partial or complete occlusion of their lumina by secondary thrombosis. A dense infiltrate of polymorphonuclear leukocytes lymphocytes and histocytes is present in the walls of the vessels and in the perivascular areas. The center of the node may show necrosis Extravasation of erythrocytes is absent (Corns) Mosinger and Jouve)

The Janeway lesions resemble the petechiae in their histologic appearance except that extravasation of erythrocytes is less evident

MENINGOCOCCEMIA (MATERHOUSE FRIDERICHSEN SINDROME

In fulminating septicemic infections with Neisseria meningitidis furpura is common. Death may occur within from 12 to 24 hours with few or no signs of meningeal involvement. On automs, home tharee

a couse ruderichsen syndrome occurs in septicemic infections with organisms other than Aeisseria meningi tidis such as Strej tococcus hemolyticus and Pseudomonas aeruginosa B Proconeus)

Ma

Histopathology The purpura of meningococcemia is caused by degeneration and inflammatory invasion of the walls of blood vessels vascultus) (see page 127) The vessels are dilated engorged with blood and frequently thrombosed. The endothelial cells are swollen and desquamating and the vessel walls show necrosis Polymorphonu

lumina of apocrine glands. The infection extends through the subcutaneous fat by way of the lymphatics to other apocrine glands is well as to eccrine glands. The lymph ressels are distended and contain many leukocytes and clumps of cocci. In the early strag, the infiltrate is composed predominantly of neutrophils. Later on, lymphocytes and plasma cells predominate and foreign body giant cells may be present. The upper parts of the dermis and the epidermis are not involved until extensive destruction has occurred throughout the subcuts.

PYODERMA GANGRI NOSUM (CHRONIC UNDFRMINING BURROWING ULCER)

Pyoderma gangrenosum begins with cutaneous abscesses which break down, forming ulcers. The ulcers spread peripherally. The advancing border of the ulcers is purplish red, edematous and under mined. The condition is associated not infrequently with ulcerative column.

Histopathology. The histologic appearance is not diagnostic. In the region of the ulcer, the epidermis is absent. The upper dermis shows necrosis and is permeated by an acute inflammatory infiltrate. Farther down, the infiltrate is chronic inflammatory, granulomatous in nature, consisting of lymphocytes, neutrophils, plasma cells his nocytes and fibroblasts. Tpithelioid cells and foreign body grant cells.

The infiltrate may extend deeply into the subcutaneous layer in areas of healing, fibrosis occurs

The epidermis at the edge of the ulcer often shows considerable proliferation, so that the histologic picture of pseudo-epithelioma tous hyperplasia (see page 334) may result. Intra epithelial abscesses may occur in this region.

Differential Diagnosis. Any ulcer, whatever its genesis may pre sent the histologic picture just described. If the biopsy specimen is taken from an area in which the epidermis shows marked hyperplasia differentiation from bromoderina may be impossible, since bromoderma too shows marked epithelial hyperplasia and intra epithelial abscesses

SUBACUTE BACTERIAI INDOCARDITIS

Three types of cutaneous lesions may occur in subrcute bacterial endocarditis petechiae, Osler nodes and Janeway lesions (Libman and Friedberg)

In old lesions, no pustulation remains and the greatly thickened horn) layer consists largely of fully keratinized cells with only a few areas of parakeratosis (Herold and Smith)



Fig. 7: Kerstons biennorrhagua late stage. Los. magnification There a ver which, paraketatothe chron there (b) permeated by numerous neutrophils. The upper stratum mylughus site seat of a spongatorm pistude (S.P.) The spongatorm appearance is caused by the preservation of the cellular membranes of epidermal cells within the pustule There are elongation of the rete ridges and edema of the psyllate. (A)D.

Connecocci can be demonstrated only rarely in el a l

lesions in 5. Only one case is on record in which gonococci were recovered on direct culture of the kissons (Margolin). It is likely that many of the cases reported in the literature as keratosis blennor clear leukocytes are found in the dimaged vasculir walls as well as perivascularly Large and small areas of hemoirhage are present in the tissue.

Meningococci can be demonstrated in the cytoplasm of endothelial cells and polymorphonuclear leukocytes. They are also found free in the lumina of vessels and in the perivascular spaces. In addition the thrombi frequently contain meningococci. Although these or ganisms can be recognized in routine stains, they are best demonstrated by Gierma stain.

KERATOSIS BLENNORRHAGICA (GONORRHEAL KERATOSIS)

Keratosis blennorrhagica occurs in chronic gonorrhea usually in association with urethral discharge and polyarthritis. The lesions have a predilection for the pilms, the soles, and the glans penis. Early lesions are represented by pustules. Graduilly, the lesions become covered with thick, horny crusts. Confluence of neighboring lesions leads to the formation of extensive horny excrescences, which have been compared with mountain ranges on a relief map.

Renter's disease, which in typical cases consists of the trind of urethritis, arthritis and conjunctivitis, may present cutaneous lesions identical with those of keratosis blennorrhagica, but gonorrhea is not the cause of the disease

Histopathology. The first histologic changes consist of the appear ance of an acute inflammatory infiltrate in the dermis and the formation of pustules in the uppermost epidermis (Carr and Fried man, Epstein and Chambers). As a rule, the pustules are spongform (Fig. 77) (Kogoj) as in acrodermatius continua and in impetigo her petiformis (see pages 102 and 104). The spongiform appearance of the pustules is caused by the presence of neutrophils inside of edematous degenerated epidermal cells, whose cellular membranes traverse one pustule like the network of a sponge (Fig. 78).

Simultaneously with or shortly after the formation of the pustules the rete ridges become elongated. It this stage, the histologic picture is indistinguishable from that of accodermatics continua-

However, as the lesions age the parakeratotic horny layer thickens to a degree that is not observed in acrodeimatitis continua and is typical of keratosis blennorrhigua. The greatly thickened horny layer is the anatomic substrate for the mountain relief appearance of the lesions observed clinically. The horny layer may measure see real millimeters in thickness. It consists of parakeratotic cells intermingled with neutrophils (Fig. 77)

In old lesions no pustulation remains and the greatly thickened horny layer consists largely of fully keratinized cells with only a few treas of parakeratosis (Herold and Smith)



in in terators blemowthagera lare stage. Los magnificitis in Tiere is a very third parakeratine horns layer (H.L.) permeased by numerous neutrophils. The upper stratum indipplin is the sert of a springform pustule (8P). The spongform appearance is caused by the presentation of the cellular membranes of epidermal cells within the pustule. There are elongation of the receivings and edema of the papillae (x100).

Gonococci cin be demonstrated only rarely in the lesions of kera tous blennorthagina. Meale and Singletary found that out of a total oil 11s case reported in the literature. Grain negative diplococci were demonstrated in tissue sections in 5 cases and in simeras from the fessions in 5. Only one case is on record in which gonococci were recovered on direct culture of the Jessons (Margolin). It is likely that many of the cases reported in the literature as keratosis blennor.

rhagica were nongonorrheal and represented instances of Reners disease (Kuske)

In REITHR'S DISLASE the histologic picture is the same as in kera tosis blennorrhagica (Lever and Granford)



1 ic 78 Keratosis blennorrhagica Tite stage High magnification of Figure 77 The spongiform nature of the pustule is well apparent (×490)

TUBERCULOSIS

When a normal not previously infected times pig is inoculated intracutaneously with an idequate dose of tubercle brillia a hard nodule develops at the site of inoculation after from 8 to 12 days. The nodule soon ulcerates. The regional lymph nodes become en larged and sometimes ulcerate (Ghon complex). Histologic examination of the primary ulcer from 10 to 14 days after inoculation reveals a marked inflammatory response with many polymorphonuclear leukocytes and tubercle bactli. During the next 2 weeks the histologic picture changes. Lymphocytes and epitheloid cells appear and replace the polymorphonuclear leukocytes. Distinct tubercles or tuberculoid structures develop at the site of inoculation, and in the regional lymph nodes as well, within 3 to 4 weeks after the moculation. Simultaneously with the appearance of epitheloid cells the number of tubercle bacilii decreases rapidly (Sulzberger).

Appeal tubercle consists of an accumulation of entitlelioid cells surrounded by a wall of lymphocytes. Usually a few Langhans giant cells are present among the epithelioid cells. The epithelioid cells center of the tubercle may show various degrees of causation. If such typical tubercles are present, one speaks of a tuberculous infiltrate Frequently, however, in tuberculous one does not find typical tubercles but only irregular accumulations of epithelioid cells within an inflammatory infiltrate, with or without caseation, and with or without Langhans giant cells. In that case, one speaks of a tuberculoud infiltrate.

It is important to realise that a tuberculous, and particularly a tuberculoid infiltrate does not necessarily mean tuberculois. Either may occur in many other diseases, particularly syphilis, leprosy and several of the deep fungus infections. The Jadassohin Lewandowsky law states that, wherever micro organisms or their products are being overcome or neutralized by the local immunobiologic reactions, tubercles of tuberculoid structures have a tendency to appear

Of interest is the work of Sabin, who found, by intraculary ous injections of various fractions of tubercle breilli, that theybrotein fraction evokes the nectorizing and lymphocytic response and the aphospholipid fraction evokes the printelloid cell response

The histologic diagnosis of the various types of cutaneous tuberculosis is dependent on the correlation of the degree of caseation necrosis the amount of inflammatory infiltrate, the degree of vascular changes and the relationship of the tubercles to blood and lymph vessels.

Assauson necrous us so called because of the cheesy macroscopic appearance of the affected ususe Microscopically, areas of cascation necrosis show <u>complete loss of liber architectural outline</u> One observes cosmophilic granular material in which unless the caserinon necrosis is far advanced some nuclei are still present Phonever most of the nuclei show palmosse-(shrunkage) or karyorthexis (fragmentation) in local tuberculous infections, cascation necrosis) is caused by the action of bacterial towns alone in hematogenous infections, oblitetatise changes in the vessels are responsible in part If any marked inflammatory infiltrate is absent in lessons of tuberculous, it is indicative of either a relative resistance of the host or of an attenuated mifection

No generally accepted classification of tuberculosis of the skin exists. The classification presented in Table 3 is based on the classifications offered by Montgomery and by Laymon and Mithelson. 176

TABLE 3 -CLASSIFICATION OF TUBERCULOSIS OF THE SAIN

	Amount of Caseation
1 Primary tuberculosis	
Localized infection tuberculous chancre	Considerable
Whem to be the skin to the	Considerable
2 Reinfection tuberculosis	ì
a Localized infection	
O(1) Lupus vulgaris	Slight
(2) Tuberculosis verrucosa cutis	Moderate
√(3) Scrofuloderma	Considerable
▼(4) Tuberculosis cutis orificialis	Considerable
b Hematogenous disseminate infection tuberculids	1
(1) Micropapular tuberculid either purely papular or rosa cea like (Lewandowsky)	None
Q(2) I upus miliaris disseminatus faciei	Slight
(3) Papulonecrotic tuberculid	Considerable
6 (4) I ichen scrofulosorum	None
(5) Frythema induratum	Considerable

1 PRIMARY TUBERCULOSIS

Primary infection with tuberculosis usually takes place in child lood. It occurs only very rarely on the skin. In the vast majority of cases, it presents itself in the lung as the so-called Ghon complex or primary complex. This consists of a small, caseous lesion at the periphery of one lung with caseation of the regional hilar lymph nodes. The Ghon complex does not become chronic. It either heals or extends rapidly. Extension may be by continuity or by hematogenous dissemination. In the latter case, miliary tuberculosis may develop.

A TUBERCULOUS CHANCRE

Primary infection of the skin with tuberculosis is more apt to occur in children but may be seen in adults (Michelson, 1935). The cutane ous lesion usually consists of a crust covered ulcer. It is referred to as tuberculous chance or primary inoculation tuberculosis of the skin. The regional lymph nodes are enlarged and may or may not suppurate and produce draining sinuses.

Histopathology. The histologic development of the lesion is very much like that observed in experimental cutaneous inoculation of the guinea pig with tubercle bacilli (see page 174). During the early

and it ad core the histologic picture is that of a handle acute,

necross After from 3 to 6 weeks, a more specific instances and develops. Epithelioid and Langhans grant cells are then present, though typical tubercles do not form, as a rule Cascatum-necross remains a prominent feature. At this stage, the number of tubercle bacult is so greatly decreased this it may be impossible to demon strate them in histologic sections and the only proof of their presence is through positive animal inoculation experiments. Simultaneously, with the decrease in the number of tubercle bacult in the lesion, the tuberculin test, pro-south negative, becomes positive. The histologic picture in theyer joinal lyimph nodes is identical with that of the cutaneous lesion (O Leary and Harrison).

R CENERALIZED MILIARY TUBERCULOSIS OF THE SMIN

The cutaneous lessons are usually papules but may be vesicles or pustules. They tend to break down and form small ulcers

Histopathology. In early lesions, one observes a nonspecific in flammatory infiltrate with loc of necrosis but without giberculoid reaction. Numerous tubercle bacilit are found within the blood vessels and in the foci of necrosis. At a later stage, juberculoid formations may be encountered (Wise).

2 Reinfection Tuberculosis

The immunity acquired by the primary infection almost always protects at least for several years. After such a latent period crinfection may occur. Reinfection tuberculosis usually is mono-organic, so that in cases in which the skin is affected other organs are, as a rule, free from active tuberculosis. As in primary tuberculosis, the infection of the skin may be localized or disseminate.

A LOCALIZED INFECTION

In localized infections one may find, depending on the virulence of the bacilli and the degree of resistance of the host, either a slight amount of taseation, as in lupus sulgars, a moderate amount of caseation as in tuberculosis verticosa cutts, or a considerable amount of caseation as in scrofuloderma and tuberculosis cutts orthicals.

(1) Lupus l'ulgaris

In lupus vulgaris the lessons, which are found most commonly on the dace consist of sharply demarcated, reddish brown patches con taining pinhead sized, deep-seated, nodules) If the blood is pressed

out of the skin by pressure with a glass slide (diascopy) the nodules stand out clearly as yellowish brown macules. Because of their yel lowesh brown color, the nodules are referred to a apple gelly nodules. In the course of time as a rule, the affected areas become atroping with contraction of the tissue. However, some areas may show yer.



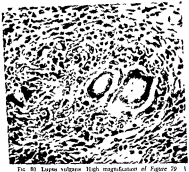
Fig. 79 Lupus vulgaris Low magnification. There are several tubercles. The large tubercle in the center shows slight cascation necrosis. (×100)

rucous thickening (lupus vulgaris verrucosus) or superficial ulceration. Squamous cell carcinoma may develop at the margin of the ulcers.

Histophilology Typical tubercles with epithelioid cells grant cells and a peripheral zone of lymphocytes are present [Caseunon necrosis within the tubercles is slight and may be absent [Figs 79 and 80]. The amount of inflammatory infiltrate composed of lymphocytes and plasma cells varies. In some cases, the inflammatory infiltrate dominates the histologic picture so that one has to search for occasional tubercles, in other cases, it is slight.

The infiltrate of lupus vulgaris is most pronounced in the upper dermis but in some areas may extend into the subcutaneous layer it causes destruction of the cutaneous appendages. In areas of healing, extensive fibrous occurs

Secondary changes in the epiderinis are common in some areas the epiderinis may show acanthosis hyperterausis and even papil



tubercle containing several Langhams grant c lls is shown. The nuclei of the grant cells he in thorseshoe arrangement. (X400)

lamatons (lupus sulgaris vertucosus). In other areas, by the pressure of the infiltrate atrophy and even destruction of the epidermis may occur resulting in ulceration and secondary progenic infection. At the margin of such inters, pseudo-epideliomatous hyperplasta and, in some instances, squamous-cell carcinoma may be found.

Tubercle bacill are present in such small numbers that their presence can hardly ever be demonstrated by framing methods. Guinea Ing inoculation on the other hand, usually gives a positive result

Differential Diagnosis Differentiation from of coidosis may be very difficult and occasionally is impossible. Vo absolute histologic criterion exists by which the two diseases can be differentiated with certainty. As a rule, however, the infiltrate in sarcoidosis tends to

JΩΛ

he in scattered islands throughout the dermis, while in lupus val garis the infiltrate tends to be compact and to be munly in the upper dermis Turthermore surcoidosis usually shows much lesslyimpho cytic reaction more fibrosis around the nests of epithelioid cells a smaller number of grant cells and complete absence of necrosis (Ron chese) The epidermis in surcoidosis is either normal or shows pres sure strophy while in lupus vulgaris in addition to atrophy there may be areas of ulceration acanthosis and pseudo-epitheliomatous hyperplusic However the only laboratory procedure by which the two diseases can be differentiated with certainty is guinea pig inocu lation which usually is positive in lupus vulgaris always negative in sarcondosis

For a discussion of swimming pool granuloma, which is thought by some to be an inoculation lupus vulgaris see page 142

(2) I uberculosis I errucosa Cutis (Including I erruca Necrogenica)

This form of tuberculosis represents an infection of a nearly im mune skin with virulent tubercle bacilly. The lesions of which there may be one or several consist of verrucous hyperkeratotic areas sur rounded by an inflammatory border Crusts may be intermingled with the keratotic material Frequently pus may be expressed from fissures within the verrucous lesion

Histopathology The histologic picture shows aganthosis hyper Leratosis and papillomatosis of the epidermis Beneath the epidermis there is an acute inflammatory infiltrate of polymorphonuclear leuko extes and lymphocytes with abscess formation. In the middle portion of the dermis one usually finds typical tubercles with a moderate amount of casertion necrosis At times however only a nonspecific inflammatory infiltrate is present. Tubercle bacilli are more numer ous than in lupus vulgaris and therefore occasionally can be demon strated histologically (Montgomers)

(3) Scrofuloderma (Tuberculosis Cutis Colliquatica)

Scrofuloderma may originate in the subcutaneous tissue fre Denomination may originate in the subcutaneous tissue frequently however it represents a direct or lymphatic extension to the skin of an underlying tuberculous focus located usually in a lymph node or bone. The lesion becomes first maintest as a bluish red printess swelling, which suppurates and later breaks down to form an ulcer with irregular undermined bluish borders. Histopathology The center of the lesion shows nonspecific abscess formation with ulceration of the epidermis. At the periphery how

ever one sees tubercle formation with marked <u>Cascation necessis</u> and a considerable amount of <u>chronic inflammators</u>, reaction (Fig. 81). The number of tubercle bacilli is rusually sufficient to enable one to find them in sections stained after Ziehl Neelsen (Michelson, 1924).

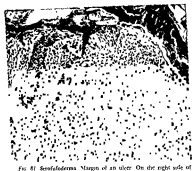


Fig. 81 Scrofuloderma Margin of an ulcer. On the right side of the photograph one observes nectors of epidermis and dermis. In the center are tuberculoid structures. On the left, the infiltrate is composed of lymphocytes and plasma cells (X200).

Differential Diagnosis For differentiation from crytheina induratum see page 186 and from gummatous syphilis, see page 215

(4) Tuberculosis Cutis Orificialis

The lesions are shallow ulcers with a granulating base, occurring about the mucous orifices of patients with virulent internal inheritations.

Histopathology The histologic picture may show merely an ulcer surrounded by a monapectife influmning millerie. In most in stance, however one finds juberel formations with egustation, necessarily the production of the ulcer than the production of the ulcer may show hyperplasta. Tuberelelpicili usually can be demonstrated in the sections even when the histologic appearance is nonspectife.

B HEMATOGENOUS DISSEMINATE INFECTION TUBERCULIDS

Tuberculids are caused by a hemitogenous dissemination of tubercle bacilli in patients possessing a rather high immunity to tuber culosis. Becuise of the high immunity, the bacilli are rapidly destroyed in the skin and therefore are usually not demonstrable in sections or by animal moculation. Although, naturally the hematogenous dissemination must take place from antimernal focus often this focus cannot be found. Clinically as well as histologically, transitions between the various forms of tuberculids occur not infrequently.

(1) Micropapular Tuberculid (Including Rosacea like Tuberculid of Leu andousky)

The eruption is limited to the facefrind consists of numerous small slightly indurated but not elevated papules of about pinhead size



Fig. 82. Micropapular tuberculid (rosacea like tuber culid of Lewandowsky) In the center there is an island of epithelioid cells with only a slight admixture of lym phocytes. This picture is indistinguishable from that of sarcoid (X100)

Some of the patients present in addition a diffuse erythema of the face To these cases the term rosacea like tuberclid of Lewandonsky

has been applied

Histopathology Histologically the papules usually show islands of epithelioid cells with only a few temphocytes and few or notgrant cells they thus present a picture indistinguishable from that of sar coidosis (Fig 82) Alaymon and Michelson) In other cases a moder ate admixture of lymphocytes is present so that the appearance re sembles lupus vulgatis more than sarcoid (Mackee and Sulzbetger) Caseation necrosis is absent as a rule occasionally one finds slight central necrosis in the tubercles (Wile and Grauer)

Since the papular type of some rosacea may show the same his tologic picture as the rosacea like tuberculid of Lewandowsky (Miescher Laymon) (see page 124) the latter diagnosis should never be made on the basis of the histologic findings alone but only when the following additional criteria exist (1) presence chinically of minute lupoid nodules and (2) concomitant evidence of rubercu losis such as pulmonary tuberculosis or a high degree of tuberculin sensitivity (Laymon). The possibility exists that all cases of rosacer like tuberculid are instances of acne rosacea, and that cases of micro papular suberculid without rosaces like features represent miliary sarcoidosis

(2) Lupus Miliaris Disseminatus Facier

Lupus miliaris disseminatus faciei occurs on the face only. The cruption consists of firm elevated discrete papilles occurring singly or in groups

o tipus vuisatis although in the latter caseation is itss pronounced

I RHENOR TERFECTION A variant of lupus miliaris disseminatus factes with the same histologic picture has been described by Ockuly and Monegomers under the name lichenoid tuberculid There is a generalized eruption of discrete or grouped erythematous papules with predominance on the extremities and usually no lesions on the face

(3) Papulonecrotic Tuberculid

The exuption is not limited to the face as it is in the two preceding types In addition to the face the extremities and the trunk may be iffected The lesions consist of indolent inflammatory papules which

come in crops and undergo central necrosis. Papulonecrotic tuber culid occasionally occurs simultaneously with enathema induration.

Histopathology. One observes a small central area of recrosis in

Histopathology One observes a small central area of accross in volving the superficial dermis and the overlying epidermis. The area



Fig. 83 Lupus miliaris disseminatus factei. A tuberot showing central cascation is present in the dermis. It is surrounded by a moderately severe inflammatory infiltrate (x100)

of necrosis is surrounded by a zone of inflammation which is largely nonspecific but may contain at its peripher, tubercles showing a mod erate amount of cascation no frosis. The infiltrate may extend into the subcutaneous fat. The blood vessels in the lower dermis show obliterative-endarteritis and endophlebitis with thrombosis and recanalization of their lumina. The wills of the vessels may be invaded by inflammatory cells. The vascular lesions are responsible for the

superficial necrosis. The histologic picture may be called a miniature erythema induratum

(4) Lichen Scrofulosorum

The lesions which occur chiefly on the trunk consist of indolent pinhead sized papules. Their color varies from that of normal skin to a pile red

Histopathology The infiltrate consists of almost pure epithelioid tubercles with only an occasional grant cell. A narrow zone of lym Ophocytes may or may not be present at the periphery of the tubercles There is no caseautor. The infiltrate is located in the upper dermis and frequently but not invariably is arranged about hair follicles (Montgomery).

(5) Ersthema Induratum

Erythema induratum or tuberculosis cuits indurativa is a chronic recuirent eruption occurring on the cales of women. In contrast with the other tuberculids the presence of tubercle pacific has been demonstrated repeatedly in histologic sections and by inoculation of aguinea pigs (Montgomery O Leary and Barker) The desions consist at first of prunless deep-seated subcutaneous infiltrations Grad ually the infiltrations eviend to the surface forming bluish red

ι.

s that y tunerations at least in some areas. Sometimes however it is necessary to cut deeper into the block of tissue to find such areas.

In area in which the infiltrate is Yuberculoid one finds epithelinedand grant cells occasionally in tubercle arrangement (Fig 85) In urers of violotypecific infiltration one observes predominantly lym, phieries and plasma cells both types of infiltrates invade between the fat cells and gradually phapes them (proliferation atrophs or Wucheratrophie of fat) Cascation necrosis in nearly always present and may be extensive in areas of cascation necrosis the fat cells often are still preserved while the invading infiltrate between them has been supplianted by an amorphous finely granular cosmophilic material in which some pyknotic nuclei are present

Ascular changes are extensive and usually severe All sizes of ves sels show proliferative changes. The changes are severest in the larger arteries and veins. Their walls become infiltrated with round cells and greatly thickened in all roats. Thrombosis and obliteration results.

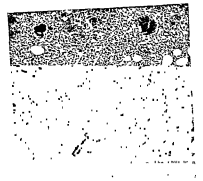
(Fig 86) The obliteration results in widespread necrosis and abscess formation. The necrosis may extend to the dermis and the epidermis and lead to ulceration.

Differential Diagnosis Differentiation from erythemyhodosum rarely cruses difficulties even though a few small foci of tuberculoid

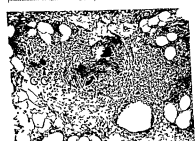


Fig. 84 Erythema induratum Low magnification. The infiltrate is limited to the subcutaneous fat. It invides between the fit cells and gradually replaces them (proliferation atroph) of the fat) (×25).

infiltrate may occur in crythema nodosim. In the first place, the infiltrate is much more[massive in crythema induration than in crythema nodosim, where it usually consists of small scattered aggregates and most important one usually finds at least a few areas of caseation[necrosis which never occurs in crythema nodosim. Further more, extensive tuberculoid/infiltration and abscess formation do not occur in crythema nodosium either and if present establish the diagnosis of crythema induration beyond any doubt. Lesions showing a pronounced tuberculoid infiltrate, extensive caseation and ulcera



proliferation of their walls (X=00)



Fic 86 Erythema induratum High magnification A large sessel (t) in the subortaneous fat is invaded by inflammatory cells and thrombosed. Vit the right of the sessel the inflammatory inflittate is nonspecific at the left epithelioid cells and a Langhans type of giant cell (GC) are located (X200).

(Fig. 86) The obliteration results in widespread necrosis and abscess formation. The necrosis may extend to the dermis and the epidermis and lead to ulceration.

Differential Diagnosis Differentiation from crythem shodosum rarely causes difficulties even though a few small foci of tuberculoid



Fig. 84 Erythema induratum Low magnification. The infiltrate is limited to the subcutaneous fat. It invades between the fit cells and gradually replaces them (proliferation atrophy of the fit) (X25)

infiltrate may occur in crythema nodosum. In the first place, the in filtrate is much more[massive in crythema induration than in cry thema nodosum, where it usually consists of small scattered aggregates, and, most important one usually finds at least a few areas of caseauon[necrosis which never occurs in crythema nodosum. Further more, extensive tuberculoid/infiltration and abscess formation do not occur in crythema nodosum either and if present, establish the diagnosis of crythema induration beyond any doubt. Lesions showing a pronounced tuberculoid infiltrate extensive caseation and ulcera

the center of the epithelioid cell islands. If a resiculum stain is employed, one sees a <u>regululum_neurond-surrounding</u> and permeating the islands of epithelioid cells (Fig. 89). The network varies in den sity. In most areas, the resiculum fibers surround almost every epithe lioid cell, but, in others, the fibers are concentrated at the periphery, leaving the central portions relatively free.



There are large islands of epithelioid cells with only a slight admixture of lymphocytes (x50)

In the Mealing phase, one for 1 recording at the petrphery of the sid the hold cells and the ly gradual transformation of reticulum fibers into collago. (* 55 can be *) recording to collago.

188

tion may resemble scrofuloderma. However, scrofuloderma shows no significant viscular changes and usually shows many tubercle bacilli on strining with /iehl Neelsen's strin

For differentiation from gummatous syphilis see page 215

SARCOIDOSIS

Sarcoidosis is a systemic disease which may affect many organs. The skin frequently is involved. Four types of cutaneous sarcoidosis are generally recognized Boeck's sarcoid subcutaneous sarcoid of Danier Roussy, lupus permo of Besnier and erythrodermic sarcoid Boecks sarcoid the most common type of cutaneous sarcoidosis is charac terized by soft brownish red intracutaneous papules nodules or plaques By central clearing annular lesions may result The lesions only rurely ulcerate Subcutaneous sarcoid of Warrer Roussy shows subcutaneous nodules which often reach several centimeters in di ameter and do not break down The overlying skin is either of normal color or bluish red Yupus permo of Besnier presents soft nodules and infiltrated plaques of violaceous color, the surface of which shows numerous telangiectases Frythrodermic sarcoid is characterized by large sharply demarcated slightly scaling brownish red patches showing little or no infiltration (Lever and Freiman Wigle) and Musso) It might be pointed out that spiegler Fendt sarcoid is not Sarcoidosis but a lymphoid hyperplasia possibly related to lymphoma (see page 491)

Histopathology The histologic picture is essentially the same in all four types of cutaneous sarcoidosis. In the skin as in other organs it is characterized by the presence of circumscribed islands of epithelioid

cells so called epithelioid cell tubercles

In Boock's sarcoid the islands of epithelioid cells are located pre dominantly in the dermis while in Darier Roussy's surcoid they are found mainly in the subcutaneous tissue. In lapus pernio the histo logic picture differs from that of Boeck's sarcoid only by the pres ence of greatly dilated capillaries in the upper portion of the dermis In the crythrodermic form the infiltrate shows rather small foci of epithelioid cells intermingled with histocytes and lymphocytes in superficial permascular arrangement (Lever and Ireiman Wigley and Musso)

In activations of sarcoidosis the islands of epithelioid cells con trun few Langhans giant cells or none at all (Fig. 87) A slight and is present particu

(Fig 88) Caseation

necrosis is almost always absent Raiery some necrosis is found in

Schaumann, 1936) Numerous orguns may be involved (Thomas, Ricker and Clark, I opgeope and Freiman) The simph nodes are frequently, and the consils occisionally, involved In the sungs, foci of sarcoidosis are often present and may, on roentgenologic examination, reveal a picture resembling that of miliary tuberculosis or Hodgkins disease (McCort, Wood, Hamilton and Ehrlich). The

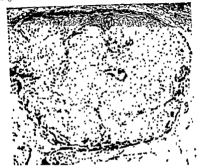


Fig. 89 Surcoidosis active stage Foots reticulum stain Reticulum fibers surround and permeate the epithelioid-cell islands. At the margin of the lesions one can observe the gradual transformation of reticulum fibers into collagen fibers. (x100)

office may be considerably enlarged The phalanges of the fingers and the toes may show lessons of sarcoidosis, which in roentgeno grams look like cysts and formerly were called ostutis cystica of Jung ling Histologically, these lessons represent replacement of bone mirrow and bone by sarcoid tissue (Ellis Weoparoutis of Heer lordt is also a manifestation of sarcoidosis (Pinner, Michelson). In occasional instances, the fleatit, the kidneys, the gastro-nitestinal tract and the central nervous system are the site of lessons (Longcope and Frenman). Involvement of the lungs, if extensive, of the heart or of the central nervous system may cause death.

Cause The cause of sarcoidosis is not yet known Some authors regard it as a noncaseating form of tuberculosis (Pinner). A few nated and often calcified (Schrumann 1941) In other cases asteroid inclusion bodies have been found inside of grant cells (Fig 91) (Lever and Freiman). The significance of the Schrumann and the asteroid inclusion bodies is not known. Neither is pecific for sarcoidosis since Schaumann bodies may occur also in berylhosis (see page 141) and insteroid bodies have been observed in other granulomas especially in foreign body granuloms.



Fig. 88 Streoidosis active stage. High mignification of Figure 87. The island of epithelioid cells is sharply de mircated. A slight lymphocytic infiltrate is present at the margin of the island. Giant cells are absent (x200).

Differential Diagnoy's The differentiation of lesions of sarcoidosis from those of lupus vulgaris may be very difficult in cases in which the islands show more than just a slight admixture of lymphocytes (see page 179). A histologic picture identical with that seen in sarcoidosis may occur in the cataneous granulomas of systemic berylliosis (see page 141) and in thoerculoid leprosy (see page 195). In the latter honever, epithelioid cell islands usurily are found invading and de stroying nerves in the dermis and the subcutaneous tissue.

Systemic Lesions The systemic nature of sarcoidosis at first not recognized, has been established particularly by Schaumann's work

authors have reported the finding of tubercle bacilli fin lesions of sarcoidosis (Kyrle kalkoff and Mohr). It is possible however that in such cases either the diagnosis of sucoidosis was erroneous or a concomitant tuberculosis evited.

If sarcoidous is due to tuberculosis it is necessary to assume that the tubercle breilit are quickly killed and disintegrated under the influence of local immune forces. Under these circumstances the lipids of the bacillus might be capable of inducing the epithelioid cell response whereis the protein of the bacillus in the absence of hypersenitivity produces no significant necrosis or constitutional reaction (see page 175). Thus the same immunologic situation responsible for the prompt killing of the bacillus inglit explain the morphologic character of the lesion (Freiman)

LEPROSY

Leprosy is caused by the lepra bacillus Mycobacterium leprae an acid last organism

and last organism. Three types of leprosy exist lepromitous leprosy, tuberculoid (or neural) leprosy and indeterminate leprosy (Canizares Arnold 1949). The lepromatous and the tuberculoid types are definite clinical pathologic breteriologic and immunologic forms of leprosy. They do not occur together in the same patient and transformation of the fepromatous type into the tuberculoid type or tice versa is very rare (indeterminate leprosy represents a transitional step. It may teman indeterminate leprosy represents a transitional step. It may be poor prognosis for arrest or long time and then heal or may develop into fepromatous or tuberculoid leprosy. Lepromatous leprosy has a good prognosis for arrest or recovery, bulle tuberculoid leprosy the lepromatic terminate because of absence of immunity in tuber culoid leprosy it is positive in indeterminate leprosy the lepromatic in the programment of the positive or negative. Cases of indeterminate leprosy showing a positive test will probably remain indeterminate leprosy the probably remain indeterminate or will develop tuberculoid lesions those showing a negative leprominities.

the cutations of the skin cilled lepromas. These granulomas occur not only in the skin but also in the nucous maintaines of the upper respiratory tract the eyes the testes the superficially to tied nerves and the reticulo-endoticiall structures such as the humps nodes the hear the sphere and the bone marrow Neurologic changes such as angesthesia trophic disturbances and partissis unally are present. Witherculoid leprosy also called neural or mixello anesthetic leprosy affects mainly the skin and the nerves

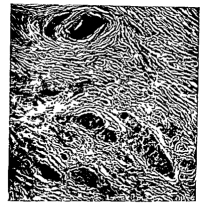


Fig 90 Sarcoidosis healing stage There is considerable fibrosis with progressive obliteration of the epithelioid cells In contrist with early lesions giant cells are numerous (x200)

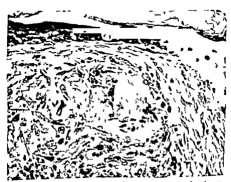


Fig 91 Sarcoidosis A large giant cell contains an asteroid inclusion, body (×400)

authors have reported the finding of tubercle bacilli fin lesions of sarcoidous (kyrle kalkoff and Mohr). It is possible however that in such cases either the diagnosis of sarcoidous was erroneous or a concomitant suberculosis existed

If sarcoidosis is due to tuberculosis it is necessary to assume that the tubercle bacilli are quickly killed and disintegrated under the influence of local immune forces. Under these circumstances, the lipids of the bacillus might be capable of inducing the epithelioid cell response whereas the protein of the bacillus in the absence of cen response whereas me protein of the bactim in the observe of hipperensimity, produces no significant necrosis or constitutional reaction (see page 175). Thus the same immunologic situated responsible for the prompt killing of the bactili might explain the morphologic character of the lesion (Freiman).

LEPROS\

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Three types of leprosy exist lepromatous leprosy tuberculoid (or neural) leprosy and indeterminate leprosy (Canizares Arnold 1949) The lepromatous and the tuberculoid types are definite clinical pathologic bacteriologic and immunologic forms of lepross. They d) not occur together in the same patient and transformation of the lepromitous type into the tuberculoid type or vice versa is very rate Indeterminate leprosy represents a transitional stage. It may remain indeterminate for a long time and then heal or may develop into leptomatous or tuberculoid leprosy Leptomatous leprosy has a not reproduced to truescentral repress Exponential repress has a poor prognosis for arrest or recovery. While tuberculoid lepross has a good prognosis for arrest or recovery. In lepromatous lepross the repromit test is negative because of absence of immunity in tuber. repromit text is regime octation or toward or minimum; in con-cided deproy it is positive in indeterminate lepros, whe lepromin text may be positive or negative. Cases of indeterminate or will develop under cut with probably remain indeterminate or will develop tuberculoid lesions those showing a negative lepromin text in all pretat t

ele

these granulomus occur not only in the skin but also in the mucous These granulomis occur not only in the same out 180 in the induced membranes of the upper respiratory tract the eyes the testes the superfactally locuted nerves and the retucio endotherial structures such as the lymph nodes the later the spicers and the bone marrow eventologic changes such as anothers trophic disturbances and paralysis usually are present. Wheretuloid leptorsy also called neural or mixedo-mesthetic leptorsy affects mainly the skin and the nerves

The cutaneous lesions consist of sharply circumscribed, erythem tous, often hypopigmented patches or fire infiltrations. Annular configuration is common. The lesions usually are hypoesthetic. The peripheral nerves, especially the ulnar nerve? are thickened and palpable. Anesthesia, troplic disturbances and paralysis occur just as

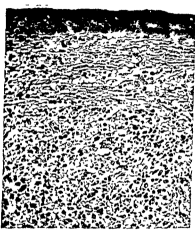


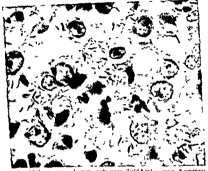
Fig. 92 Lepromatous leprosy, early stage. The granuloma tous infiltrate consists predominantly of histocytes and lepracells with formy cytoplasm (×200).

in the lepromatous type Indeterminate leprosy resembles tuberculoid leprosy in its cludeal manifestations

Histopathology Repromatous leprosy shows a granulomatous in filtrate. It is massive in the upper dermis and focal around the arteries, the veins and the nerves in the lower dermis and the subcataneous fat. Histocytes and lepra cells predominate, but, in addition there are lymphocytes, plasma cells and in older lesions, fibroblasts (Fig. 92) Lepra cells, or Virchow cells develop from histocytes. They are large, form, cells resembling antihoma cells. On staining with fat stains, they are shown to contain lipid which, in contrast to

that found in xanthoma is not doubly refractile (Tilden) On stain

stressed that Mycobacterium leprae, unlike Mycobacterium tubercu losis is not strongly acid last. Therefore sections must be decolorized



116. 93 Lepromatous let ross early stage Zield Neelsen stran Numerous letra bacilli are present. In the left center, two bundles of bacilli, are shown (x800).

phritis (Condr) (Fig. 91) On stanning after Tield Neelsen or Fire their may not contain large globi of partially degenerated lepist bacilli

Muberculoid lepros, shows a suberguloid infiltrate. Patients with suberculoid lepros, have fairly good immunity against the lepra bacillus. This explains in accordince with the Jadassohn Lewandow sky liw (see page 175), the scarcity or the obsence of floacills and the suberculoid tissue responses Sections may show almost pure engitled.

oid-cell tubercles so that differentiation from sarcoidosis may be difficult (Fig. 95). However, a thorough search usually will reveal invasion and destruction of nerves in the dermis or the subcutaneous tissue by epithelioid-cell tubercles, which never occus in sarcoidosis In some cases, the tubercles show a moderate admixture of Jimpho cytes and contain, giant cells. However, Acascation necrosis does not occur in the skin. Lepra bacilli may be absent in the lessons of tu

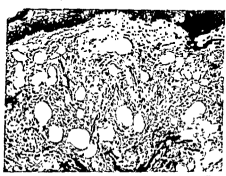


Fig. 91 Lepromatous leprosy, late stage. High magnification. There is some fibrosis. The lepra cells are very large in size and appear as round empty spaces. (X200)

berculoid leprosy, but not infrequently they are found in small numbers after prolonged search (Arnold, 1915)

Indeterminate lepiosy sho around the vessels and the nerves of ind only with difficulty (Canzares)

NTURAL LESIONS Lesions in the large peripheral nerves occur in almost every case of leprosy, regardless of type. In all three types, the histologic appearance of the neural lesions corresponds to that of the cutaneous esions (Pardo Castello, Tiant and Puñeyro). The nerve lesions of depromatous leprosy show large vacuolated Virchow cells and numerous lepra bacill. In absertational leprosy, one observes in the nerves an extensive infiltrate of epithelioid cell tubercles containing few or no bacilli. However, in contrast with the cutaneous lesionsacascation necrosis is common. In occasional instances, mas

sive caseation of the tuberculoid lesions with complete destruction of nerve tissue occurs ("Offiquative neutrus").



Fig. 95 Tuberculoid Leptory Low magnification. The militrate consusts of epithelioid-cell subercles aboung a single admicture of lymphocytes particularly at their surgins. That the histologic picture is very much like that of actiondoms (X100).

DIPHTHERIA OF THE SKIN

The clinical picture of diphilieria of the skin is varied. Most commonly encountered are one on executely geometed conclusers exercited with a pseudomembrane the removal of which is difficult and followed by profuse pherling. Next in frequency are most, crusted exerminous lesions which occur particularly in the terto autricular region Other types of lesions occasionally observed include impetigations, research and bullons (Resia).

Histopathology. The histologic picture of all forms of diphthetia is nonspecific. The ulcerative lesions show destruction of the epidermis and of the upper dermis within the region of the ulcer. At the margin of the ulcer, the epidermis is acanthotic. The dermis be neath the ulcer shows evidence of acute inflammation. The floor of the ulcer is covered with a layer composed of necrotic cells fibrin and numerous neutrophils. In some instances, numerous klebs Loeffler bacilli can be seen in this necrotic layer (Allen). These brothly are borders and tend to be in paliside like arrangement. The finding of such bacilli is suggestive of diphtheria. However, since diphtheria bacilli cannot safely, be differentiated from pseudo diphtheria bacilli in stained sections, smears and cultures are necessary for the establishment of the diagnosis.

The eczemnious lessons may show the histologic picture of either an acute, a subacute or a chronic derimitus. The histologic changes thus are identical with those observed in an ordinary bucterial dermatus (Robert).

ANTHRAX

Anthrax, caused by Bacillus anthracis, is characterized by a car buncle like lesion located usually on an exposed portion of the skin. The lesion starts as a papule, enlarges, ulcerates and then becomes covered with a black eschar Trequently, a ring of vesicles is present at the margin of the eschar Marked eightema and edema surround the lesion. Suppuration does not occur. Pain is characteristically slight or absent. Often there is regional lymphadenopathy, which may be painful.

Histopathology. At the site of the ulcer, the epidermis is destroyed and the ulcerated surface is covered with incrotic tissue. The adjoining epidermis shows spongiosis and occasionally intra-epidermid vesicle formation and is invided by neutrophils. There is marked edema of the dermis separating the bundles of collagen and loosening their fibrils. Numerous crythrocytes and neutrophils are present throughout the dermis and the subcutaneous tissue. However, abscess formation is absent. Only few histocytes can be observed. The blood vessels are dilated and their walls show diffuse degenerative changes.

Anthrax breilli are present in large numbers and can be recognized in sections stained with routine stains but are visualized best in sections stained with Gram's stain. The anthrax breillis is a large, rod shaped, square ended. Gram positive breillis, from 6 to 10 m crons long and from 1 to 2 microns thick. In the tissue, it is usually surrounded by a well defined capsule. Anthrax breilli are found in countless numbers in the necroite tissue at the surface of the ulcer. The dermis also contains numerous breilli, while the subcutaneous

tissue usually contains only a few. It is worth noting that phagocy tosis of the bacilli by either neutrophils or histocytes is absent (Lebo with Mckillip and Conboy)

TULAREMIA

Tularemy, an infectious diserse caused by Bacterium tularense, occurs in whire types the ulceroglandular the oculoglandular, the glandular and the typhoid types Specific cutaryosis lessons occur only in the ulceroglandular type. They are of two varieties ulcers and nodes. One or several ulcers occur is primary lessons at the site of infection they tond to have a <u>nunched-out appearance</u>. Cutaneous or subcutaneous odes form in the small lymph nodes located along the lymph vessels draining the primary lesson they are hard and tender at first but may proceed to suppuration.

Histopathology. The primary ulcer shows a nonspecific granular tion tissue into which are embedded granulomas composed of three innes. (I) a central zone of necrosis, consisting of finely granular cosmophilic material nuclear fragments and a few crythrocytes (2) in intermediate zone composed of epitheloid cells with a few langhans grant cells and lymphocytes and (3) an outer zone con sixing largely of lymphocytes but containing also some histocytes playing cells and extra-asared crythrocytes. Accular changes in the outer one and in the surrounding granulation tissue are conspicuous they consist of proliferation of the productional cells and militation of the vascular walls by inflummatory cells (Schuermann and Reith) Older lesions may show epitheloid-cell ubercles with only a slight inflammatory reaction and thus may have a surcoid like appearance. Bacterium tularense although prevent does not stain in the sections.

The cutaneous "Timphatic nodes show like the primary ulcer of tularemia scattered granulomas in which three zones can be recognized

Differential Diagnosis. The histologic picture differs from that of tuberculosis by the (distinct arrangement fin three zones which is rurely seen so cleritly in tuberculosis by the presence of (ascular change) and by the presence of (erridinocties in the central hiercrost-guid and the outer hymphocytic zone (Reich). Differentiation from portorichois (see page 230) and hymphogyriunloma venereum (see page 252) however may be impossible.

CHANCROID

Chancroid caused by the streptobacillus of Ducrey (Hemophilus ducrey) is a venereal disease causing one or several ulcers chiefly

in the genital region. The ulcers show little if any induration and tend to have an undermined border. Regional lymphidenitis is common and usually terminates in abscess formation (buba).

Histopathology. The chancroidal ulcer presents a granulomatous reaction which is sufficiently distinct to permit a diagnosis of chan croid in miny instances (Heyman, Beeson and Sheldon) The lesion consists of three zones (Sheldon and Heyman) and shows chiracter istic vascular changes (Pund, Greenblatt and Huie) The surface zone or base of the ulcer is rather narrow and is made up of polymorphonuclear leukocytes fibrin, red blood cells and necrotic tissue Below this is a fairly wide zone of edemitous tissue composed munly of endothelial cells in varying stages of proliferation Newly formed blood vessels are numerous. There is mirked endothelial proliferation within the vessels, frequently blocking their lumina and leading to thombosis. In addition, there is degeneration of the wills of vessels. The third zone is composed of a dense infiltration of plasmicells and lymphocytes.

Demonstration of Ducrey becille in the tissue by straining with Giemsa's stain, Gram's stain or with polychrome methylene blue is only rarely possible. They are most apt to be found as Gram negretive coccobrelli between the cells of the surface zone (Sheldon and Heyman).

GRANULOMA INGUINALI

Granuloma inguinale is a venereal disease caused by Donorama granulomatis, which is found in the lesions in the form of intracyto plasmic inclusion bodies called Donovan bodies. The taxonomic position of Donorama granulomatis is as yet undecided but it is most likely a bacterium and not a virus. It grows well on the chorio allantoic membrane of chick embryos.

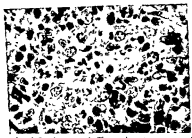
The lesions occur predominantly in the genital region and consist of shriply demarcated granulomatous areas which have a lind how and bleed easily. The border is elevated and often shows a serpiginous outline. The lesions sprend very slowly by peripheral extension and may attain large size. After years, the lesions heal with a thick, fibrous, contracted scar. In occasional instances, squimous cell carcinoma supervenes (Beerman and Sonck, Alexander and Shields).

Histopathology The epidermis may be thinned but more often shows acanthosis, which may reach the proportions of pseudo epitheliomatous hyperplasm (Beerman and Sonck) A granulomatous infiltrate is present in the dermis composed predominantly of histocytes and plasma cells Scattered throughout the otherwise mononuclear infiltrate one finds small abscesses composed of poly

morphonuclear leukocytes (Allen) The number of lymphocytes is

conspicuously small

Intracytoplasmic inclusion bodies called Donovan bodies are present within a variable number of histocytes. The parisitized histocytes or macrophages possess abundant cytoplasm and may measure 20 microns and more in diameter. Their cytoplasm has a cystic appearance. Within the cystic compartments of the cytoplasm



Fo Jf Granuloma ingu nale The granulat on tissue is composed red in nantly of I cocytes and plasma cells. There is a diffuse sprin kl g of polymorph nuclear lephocytes Several of the hist ocytes con tam D nivan bod es vith nitle r cytoplasm (×400)

one sees groups of small round or oval encapsulated bodies meas uring I to 2 microns in diameter (Fig 96) In the cross sections of large histocytes several dozen of such bodies may be observed. The Donovan bodies are recognizable in stains with hematoxylin and eosin but are seen best in sections strined with Giemsa's stain In such sections they appear bright red (Mexander and Schoch) When a silver stain is used the Donovan bodies appear black and have the shape of a closed safety pin because of their elongated ovoid shape and intense bipolar staining reaction (Torpin Greenblatt and Pund) The capsule surrounding the Donovan bodies does not stain. In addition to their intracellular location a few Done - L be to -1 .

easier to find them in tissue scrapings or tissue imprints stained with Giemsa's or Wright's strin than in fixed tissue sections

Differential Diagnosis The parasitism of histocytes is strikingly similar to that observed in rlunoscleroma histophismosis and leish maniasis However, the small size of the Donovan bodies and the presence of small abscesses in the infiltrate usually make a differen tiation from these diseases possible (See Table 5, page 237)

A difficult problem may be posed by the marked epidermal proliferation present occasionally in granuloma inguinale (Beerman and Sonck) Several biopsies may be necessary to decide whether it repre sents merely pseudo epitheliomatous hyperplasia (see page 334) or squamous cell carcinoma which occasionally supervenes in granu loma inguinale

RHINOSCLFROMA

Rhinoscleroma is a chronic infectious disease in which the nose the lip and the upper respiratory tract are infiltrated with hard plaques and masses

Histopathology. The granulomatous infiltrate is strikingly rich in plasma cells and contains two peculiar types of cells, the Mikulica cell and the Russell bodies Because of their presence, the histologic

picture of rhinoscleroma is diagnostic

The Mikulicz cell is a very large, round histocyte measuring up to 100, occasionally even 200, microns in diameter. It has a pale reticulated, ill defined cytoplasm and an eccentric nucleus. If a bac terial stain such as Giernsa's stain is used one finds within the cyto plasm of many Mikulicz cells as well as in their vicinity many breilli called Klebsiella rhinoscleromatis or Frisch bacilli (Fig 97) They are short, Gram negative rods measuring 2 by 5 microns in length and are surrounded by a narrow gelatinous capsule (called gloea) It is not certain that the disease is caused by this organism

The Russell bodies are elliptic formations, measuring from 20 to 40 microns in diameter. Thus, they are smaller than a Mikulicz cell but still twice as large as a normal plasma cell. They have a homo geneous, brilliant red light refractile cytoplasm and no nucleus (Fig 98) They form within plasma cells as a result of their degeneration

and finally are expelled (see page 37)

The amount of cellular infiltration varies with the age of the lesion Larly lesions show considerable infiltration by numerous plasma cells and lymphocytes with occasional instruction of minerous plus, many Russell bodies are present but only few Mikulicz cells Gradually, the number of Mikulicz cells increases to such extent that they predominate the histologic picture giving the section a



Fir 97 Rh noscleroma G emsa sta n There are se eral M k il α cells the cytoplasm if wl cl. s pide ret cubated and ill-defined. One M k il α cell conta ns. n its cytoplasm many Frisch bacilli which appear I era s small round deeply sta n ng bod es (X200).

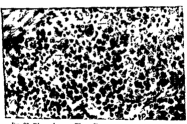


Fig. 98. Rh noscleroma. The infiltra e contrans many plasma cells and one Rus ell body. The Russell body is larger than the other cells in the infiltrate. It has a homogeneous br ll ant sed refract le cyto-plasm (x400).

easier to find them in tissue scrapings or tissue imprints stained with Giemsa's or Wright's stain than in fixed tissue sections

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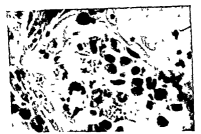
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F G 97 Rhinoscleroma Ciemsa stain There are several M kul cz cells the Gioplasm of which is pale ret cultied and ill-defined. One M kul cz cell contains in its cytoplasm many Frisch bacili which appear here as small roun! deeply sta ning bodies (X900)



Fig. 98 Rhinoscleroma. The infiltrate contains many plasma cells and one Russell body. The Russell body is larger than the other cells in the rufiltrate. In homogeneous brill ant red refractile cytoplasm. (x400).

Bacterial Diseases

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foamy, lacelike appearance Yimilla fibrosis replaces the Mikulicz cells (Kline and Brody)

Differential Diagnosis Parasitized histocytes are observed also in cutaneous leishmaniasis histoplasmosis and granuloma inguinale For their differentiation see cutaneous leishmaniasis (page 235 and Table 5 page 237) Russell bodies are not found in these three diseases and therefore are of considerable diagnostic value. However, they are not specific for rhinoscleroma because they may occur in other diseases when an infiltrate rich in plasma cells is present—for instance in sphilis tuberculosis squamous cell carcinoma and my cosis fungoides.

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Spirochetal Diseases

SYPHILIS

Acquired syphilis may be divided into three stages primary, secondary and tertuary. Primary and secondary syphilis comprise the early phase and tertuary syphilis the late phase of the infection. During the early phase, the causative organism. Treponema pallidin (spirochaeta pallida), often can be demonstrated in the cutaneous lesions by dark-field examination. In the late phase, no spirochetes can be demonstrated in the cutaneous lesions.

Primary syphilis is characterized by the syphilitic orthard chanced which is usually a single lesson but may be multiple. The typical, or Hunterian, chancre is represented by a brownish red, indurated found papule or plaque with an eroded surface. Occasionilly, the chance shows ulceration. The regional lymph nodes are enlyingd.

Secondary syphilis is characterized by a more or less generalized eruption, which is composed usually of metules or pipules having a brownish red color. In the anogenital region, the pipules may be come large, vertucous and moist, they are then called condylomata lata (They must be differentiated from condylomata acumulata, a variety of vertuca, see page 251). In some instances, the cutaneous eruption, due to the presence of scaling, resembles psoriasis (psoriasi form syphilid). Occasionally, pustules develop and are followed by heavy crusting (rupial syphilid). Ulceration of lesions in secondary syphilis is very tare and occurs only in severe cases. Alopecia syphilitica is characterized by numerous small patches of partial, "moth eaten" alopecia.

Tertiary syphilis often shows only a single lesion, but occasionally several lesions. A superficial nodular type and a deep, gummitous and

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center. Lesions of the gummatous type begin as a soil cultureous or subcutaneous swelling, which breaks down to form one or several ulces having a punched out appearance. In rare instances, just a articular nodes occur in tertiary syphilis. They are printess, slouly growing, subcutaneous, fibrous nodules, often symmetrically situated. in the vicinity of joints. The elbows and the knees are the sites of preddection

Congenital, or prenatal, syphilis shows the same cutaneous manifestations as acquired secondary and territyr syphilis with the following exceptions first occasionally bullous lesions occur especially on the palms and the soles as a minifestation of secondary syphilis in infants and second juxta articular nodes do not occur

Histopathology The fundamental pathologic changes in syphilis are (1) a predominantly perivascular infiltrate composed of lympho cytes and many plasma cells and (2) endarteritis and endophlebitis In tertiary syphilis one usually finds in addition a tuberculoid in filtrate and caseation necrosis

PRIVARY SYPHILIS In the typical primity lesion the epidermis shows at the mitgin of the lesion acanthosis. Toward the center the epidermia gradually becomes thining until in the central portion it is entirely absent (Fig. 99). An offittirate composed of lymphoties and many plasam cells is present in the dermis. It is compact in the center while at the margin it consists of individual perivascular islands. Both blood and lymph vessels show changes. The capil briefs and the lymphituses are increased in number and show considerable proliferation of their endothelial cells (Fig. 100). The larger blood vessels show proliferation of all heir coats and invision of their walls by the cellular infiltrate. Obliteration and invision of the lumin of some of the vessels is common and results in small fox. of necrosis. Occusionally, the lumin of so many vessels are occluded that massive necrosis results. In such cases, the primary lesion presents used, yet nuclear instead of an erosion.

On straining with a saint strain spirochetes usually can be found often in large number. They are found throughout the tissue but especially in and around the walls of capillaries and lymphatics.

Histologic examination of the scional lymph nodes in primary whilits may reveal only a nonspecific inflammation, not infrequently however small foci of tuberculoid reaction are encountered in addition (Michelson)

SECONDEN STITUTES In secondary syphilis the number of spirothetes seen in sections stanted with Levadria stain varies with the cape of beams. In the meeting beams to early secondary syphilis spirochetes cannot be found at a rule for spiral resions they are seen occurrently. In conditionata lata, they are also

Spirochetal Diseases

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Secondary syphilis is characterized by a more or less generalized eruption, which is composed usually of micules or pipules having a brownish red color. In the anogenital region, the pipules may be come large, vertucous and moist, they are then called condylomita lata. (They must be differentiated from condylomita acuminata a variety of vertuca, see page 251.) In some instances, the cutaneous eruption, due to the presence of scaling, resembles psoriasis (psoriasi form syphilid). Occasionally pustules develop and are followed by heavy crusting (rupial syphilid). Ulceration of lesions in secondary syphilis is very rare and occurs only in severe cases. Alopecia syphilitica is characterized by numerous small patches of partial, 'moth eaten' alopecia.

Fertuary syphulus often shows only a single lesion, but occasionally several lesions. A superficial nodular type and a deep gummitous.

- Lesions of the nodular type show an accosed of nodules and a smooth atrophic

center Lesions of the gummatous type begin as a soft cutaneous or subcutaneous swelling, which breaks down to form one or several ulcers having a punched out appearance. In rare instances, justa articular nodes occur in tertiary syphilis. They are printless, slowly growing, subcutaneous, fibrous nodules, often symmetrically situated in the vicinity of joints. The elbows and the knees are the sites of predilection

Congenital, or prenatal syphilis shows the same cutaneous manifestations as acquired secondary and tertiry; sphilis with the following exceptions first occasionally bullous lesions occur especially on the palms and the soles as a manifestation of secondary syphilis in infants and second juxta-articular nodes do not occur

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Tertury syphilis often shows only a single lesion, but occasionally several lesions. A superficial nodular type and a deep guminatous type of tertury syphilis occur. I esions of the nodular type show an active, serpiginous border composed of nodules and a smooth atrophic center. Lesions of the guminatous type begin as a soft cutrueous or subcutaneous swelling, which breaks down to form one or several ulcers having a punched out appearance. In rare instances, juxta articular nodes occur in tertiary syphilis. They are painless slowly growing, subcutaneous, fibrous nodules, often symmetrically situated.

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PRIMARY SYPHILIS In the typical primary lesion the epiderinis shows at the margin of the lesion acanthosis Toward the center the epidermis gradually becomes thinner until in the central por

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Secondary Syphilis In secondary syphilis the number of spiro chetes seen in sections stained with Levaditis stain varies with the type of lesion. In the inicular lesions of early secondary syphilis spirochetes cannot be found as a rule. In papular lesions, they are seen occasionally Incondylomata lata they are almost always pres ent in sufficient numbers to be found without difficulty. They occur not only in the dermis but also between enidermal coll

a dingn



Fig. 99 Primary syphilis I on mignification. The margin of an erosion is shown. The epidermis gradually becomes thinner as it approaches the erosion. (×100)

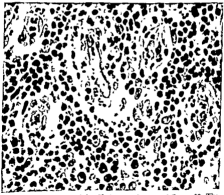


Fig 100 Primary syphilis High magnification of Figure 99 The capillaries are greatly increased in number and show marked proliferation of their endothelial cells. Many plasma cells are present in the dense infiltrate (X400)

ing of their endothelium and are surrounded by a slight infiltrate of lymphocytes and plasma cells However the number of plasma cells is not sufficiently large to be of diagnostic value

Papular lesions, as a rule, have a diagnostic appearance Not only the superficial but also the deeper vessels of the dermis are involved They show marked endothelial swelling and are surrounded by a pronounced infiltrate (Fig. 101). Because of us periciscular coat. pronounced montrate (vig tot) tocates of its best-patch, sleet elike, arrangement the infiltrate his a patchy distribution. The patchy pattern stands out very clearly in the lower dermis. In the upper dermis, in addition to the patchy infiltrate, there may be ficantly high number of

102) In the differential

ie exudative discoid and

lichenoid chronic dermatosis of Sulzberger and Garbe have to be excluded The latter in particular may suggest secondary syphilis lucause of the parchy distribution of its infiltrate and the presence of many plasma cells but neither chronic dermatitis nor exudative discord and lichenoid chronic dermatosis shows the marked endo thelial swelling of the vessels and the extension of the patchy infil trate into the lower dermis

Condylomata lata show the same changes in the dermis as the papular lesions. In addition the epidermis shows considerable acanthosis with broadening and elongation of the reje ridges. intra cellular and intercellular edema of the rete cells and migration of polymorphonuclear laukocytes through the epidermis

Psoriasiform secondary syphilis may show epidermal changes simi far to those of proriests but the derinal infiltrate is that if you -!

is a a suocornear pustules. The diagnosis of syphilis usually can be made because the lower dermis shows the characteristic in filtrate of secondary syphilis

I legrature lessons are rare in secondary syphilis. They occur when the larger vessels at the border of the dermis and the subcutaneous tissue become completely occluded by syphilitic endarteritis and endophlebitis (Wile Wieder and Warthin)

1 - 4) in many instruces however no infil trate is found (essential alopecia). Wile and Belote believe that the latter type of slove local action of

on endocrine glands or on the autonomic nervous system. Meningeal syphilis is common with that type

It is not always possible to assign on the bisis of histologic eximination, cutaneous lesions of syphilis to either secondary or ternary syphilis. Not infrequently, one finds in lesions which clinically are to be classified as late secondary syphilis an admixture of epithelioid and grant cells. On the other hand, lesions of early terrary syphilis.



Fig. 101 Secondary syphilis I on magnification. The vessels through out the detries show endothelial proliferation. The cellular infiltrate is located around the vessels in contileeve arrangement (X100).

may show the histologic infiltrate of secondary syphilis without any optibelioid and grant cell reaction

TERTIARI STIHLIS In tertiary sphills one observes a granulation tissue composed of symphosites plasma cells histocytes, fibroblasts and varying numbers of enithelioid indigiant cells. Usually but not always the number of plasma cells is prominent. The infiltrate is missive in the center but tends to have a perioriscular arrangement at the periphery Wascular changes are prominent particularly and the larger ressels. They show proliferation of their walls leading to marrowing of their liminary and invision of their walls by the inflammatory infiltrates. Assessing the viscolar changes because often the vascular changes because often the vascular changes because often the vascular changes are just as severe in areas without caseation in areas with caseation. It seems more likely that caseation represents

an allergic phenomenon paused by alteration in the reactivity of the tissue (Boyd) In the Caling stage, lesions of tertiary syphilis show numerous fibroblists. The process ends in fibrosis. Tertiary syphilis occurs in the skin in two forms nodular and gummatous

In nodular tertiary syphilis, the granulomitous process is limited to the dermis The number of sepithehold and giant cells is small,



regu magnification of Fig. ure Int The petroascular infiltrate contains many plasma cells. (×200)

is a rule. Occasionally, however, these cells are furly numerous in the center of the lesion. They tend to be without arrangement into real tubercles and grant cells of the foreign body type are more com mon than those of the Langhans type (Fig 103) Cascation Decrosis is usually not extensive and may be absent. If extensive, ulceration

In gummatous tertiary syphilis the oran 1-

usise caseation necrosis occurs in the center of the



Fig. 103 Teruary syphilis nodular type. In the center of the field a large island of epithelioid and grant cells is located. Most grant cells are of the foreign body type rather than of the Langham type. The infiltrate at the margins contains many plasma cells. (x100)



Fig. 101 Tertiary syphilis gumma. On the left side of the field, the 'm cells. In the center it Most giant cells are e sees part of the large of the eumma. (XOD)

lesson (Fig. 101). The epithelioid and the giant cells are located mainly in the vicinity of the areas of caseation. Because of the deep extension of the process not only the vessels of the dermis as in nodular tertiary syphilis but also the large vessels of the subcuta neous layer are markedly involved.

neous tayer are markenly invoices.

Differentiation of tertury syphilis from tuberculosis may be difficult. In the absence of caseation, the nodular type may resemble lupus vulgars and the gummatous type may suggest scrofuloderma or enythema induratum. The fatter in particular, may be difficult to exclude because it shows obliterative vascular changes similar to those of tertury sphilis. But aside from enythema induration, tas cultir changes provide the most important point of differentiation between suphilis and tuberculosis. In addition, the prevalence of plisma cells and the presence of guant cells of thelloreign body type rather than of the Langhans type fivor a diagnosis of syphilis.

runer than of the Langhans type I wor a diagnosis of sphillis in putsearticular nodes the histologic picture varies with the age of the lesion. Notify lesions are fairly cellular (Tuta and Coombs) and present embedded in a dense fibrous usue granulomatous areas composed of epitheliotic cells. I jumphocytes and plasma cells with an occasional Langhans type of giant cell. The blood vessels

which the histologic picture is the same as just described for early lesions an intermediate zone showing dense fibrous tissue with only few cells and an inner zone showing highinized fibrous bands rain using to form pirtutions for cystic spaces. The cystic spaces contain

jessner as well as Hu. Liu. Chen and Frizier have succeeded in producing siphilis in rabbits by inoculating them with portions of juvia articular nodes.

CONSTITAL STRILLS The Instologic changes in the cutaneous lesions of early, congenital syphilis are similar to those seen in ac quirted fecondary syphilis Except for the bullous lesions of the palms and the wise. The Fixien may show in plasma cells in the detrial infiltrate but numerous polymorphomulear leukocytes and a few lamphocytes. Obliterative vascular changes are present, however (Fraser)

Sytheris of Internal Organs It is not appropriate to discuss here in detail the histologic appearance of the lesions of syphilis in the

internal organs. However it may be pointed out that late syphilis may cause two types of reactions in the internal organs gumma and diffuse interstitual inflammation. The latter reaction is more common

The gummata in internal organs show the same histologic changes

as those observed in the skip

Diffuse interstitual syphilitic inflammation manifests itself as ac cumulations of lymphocytes and plasma cells around the small vessels Cascation is usually absent As a result of the long continued milam mation there is gradual degeneration of the parenchymatous struc tures and their replacement by fibrous tissue (Boyd) This type of reaction produces for instance syphilitic hepatic cirrhosis Syphilis of the north is also of the diffuse type. The process starts around the vasa vasorum of the adventitia. The inflammatory infiltrate then ex tends along the vasa vasorum into the media In the media one observes foci of inflammatory cells nultury necrosis and extensive destruction of elastic tissue

General paresis shows permascular accumulations of lymphocytes and plasma cells in the meninges and in the cortex of the brain. In addition there is extensive degeneration of the cortex. In contrist to general paiesis tabes does not show inflammatory changes. The essential lesion is degeneration of the posterior columns of the spiral card

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Fungus Diseases

DI RMATOPHYTOSIS (TINI A)

The following are the most important fungi crusing superficial tungus infections. Microsporum cams (lanosum). Microsporum undoutin Erichophyton neutogrophytos (gypeum). Trichophyton intrium (purpureum). Epidermophyton floccosum. Litchof hyton (Achonon). schoenleini. Malassena. Jurjur and Nocardia. (Actino moces) ministrius.

Clinically nine types of superficial fungus infections can be recognized (1) times capitis (2) times barbic (3) times corporis (4) times curity (5) dermatophytosis of the feet and the hands (6) onychomy

costs (7) fixus (8) times versicolor and (9) crythrisms

I mer capitis which occurs almost exclusively in children usually is caused either by Microsj orum causs or by Microsj orum audoum. The iffected larts tend to break off. Microsj orum causs may evoke promounced inflammation of the scalp so called kerion Celsi, while Microsj orum audoum usually produces hitle inflammatiory reaction.

Finen bribge rare in the United States is usually crusted by I richabliston mentagrot listes. It is characterized by a boggy inflant

matory infiltration in the hearded region of men

There corpores if caused by Microsporum cause manifests itself is multiple inmulti lesions with an active often vesicular border and central elevating if caused by Irichophyton mentagrophytes one lands only one or at the most a few annular lesions showing little on occurral clearing if caused by Irichophyton rubrium there are fairly sharply demarcated sheetlike areas of crythemy with slight saling

Tines crurs usually caused by I thermophyton floccosum, pro duces are sof crythiam and scaling with a slightly clevated often vestually border on the opposing surfaces of the skin between the scrotum and the thighs and in the perincil in I the glutcal regions

Dermatophytosis of the feet and the hands usually caused by trichophyton mentagrof hyter and occasionally by Pindermophyton floccosum is characterized by inaccration between and underneath

internal organs. However, it may be pointed out that late syphilis may cause two types of reactions in the internal organs, gumma and diffuse interstitual inflammation. The latter reaction is more common

The gummata in internal organs show the same histologic changes as those observed in the slin

Diffuse interstitual symbilitic inflammation manifests itself as ac cumulations of lymphocytes and plasma cells around the small ressels Cascation is usually absent. As a result of the long continued inflam mation there is gradual degeneration of the parenchymatous struc tures and their replacement by fibrous tissue (Boyd). This type of reaction produces for instance syphilitic hepatic cirrhosis Syphilis of the portrais also of the diffuse type. The process starts around the vasa vasorum of the adventitia. The inflammatory infiltrate then ex tends along the vasa vasorum into the media. In the media one observes foci of inflammatory cells miliary necrosis and extensive destruction of clastic tissue

General paresis shows perivascular accumulations of lymphocytes and plasma cells in the meninges and in the cortex of the brain In addition there is extensive degeneration of the cortex. In contrist to general parests takes does not show inflammatory changes. The essential lesion is degeneration of the posterior columns of the spiral cord

RIRI IOGR APIA

Syphilis

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Fungus Diseases

DERMATOPHYTOSIS (TINFA)

The following are the most important fungi causing superficial fungia infections. Alterosporum causi (lanosum). Microsporum audouum Trachosphyton mentagrophylise (g.pseum). Trachosphyton rubrum (purpureum). Epidermophyton floccosum, Trachophyton (Achonon). schornleim. Malassezia Jurjur and Nocardia (Actinomyces) ministisma.

Clinically nine types of superficial fungus infections can be ricrog nized (1) times capitis (2) times barbie (3) times corports (4) times cruits (5) dermatophytosis of the feet and the hands (6) onychomy

costs (7) facus (8) times versiculor and (9) erythrasina

Timea capitis which occurs almost exclusively in children usually is caused either by Microsporum earns or by Microsporum audourn. The affected hars tend to break off Microsporum earns may evoke pronounced utiliammation of the scalp, so called kerion Celsi, while Microsporum audourn usually produces little inflammatory reaction. Finea barries rate in the United States, is usually caused by

Timea barbae rate in the United States is usually caused by Truhophysion mentagrophyses. It is characterized by a boggy inflam majory inflitration in the bearded region of men.

Timea corports if caused by Microsporum cams, manifests uself as multiple annular lesions with an active often tesseular border and central clearing if caused by Freehophyton mentagraphytes, one finds only one or at the most a few annular lesions showing little or no central clearing if caused by Truchophyton rubrum, there are fairly sharply demarcated sheetlike areas of crythema with slight scaling

Tinea cruris isually caused by Epidermophyton floccosum, produces areas of crythema and scaling with a slightly elevated often vescular border on the opposing surfaces of the skin between the scretum and the thighs and in the perineal and the fluteal regions

Dermatophytosis of the feet and the hands usually caused by Inchophyton mentagrophytes and occasionally by Epidermiophyton floreosum, is characterized by maceration between and underneath the toes and an erythematous scaling and vesicular eruption on the feet and the hands especially the soles and the palms

Onychomycosis crused usually by Trichophyton rubium shows

disintegration of the nail substance

Figure 17 this country is caused by Firchophyton (Achonon) schoollem. Most commonly it affects the scalp where it produces inflammation with formation of perifollicular crusts called scuula Destruction of the hair ensues. Healing takes place with scarring

Tines versicolor caused by Malasse in furfur most commonly affects the upper trunk where one finds areas of brownish discoloration. The surface of the discolored areas shows fine branny scaling

Erythrismi crused by Nocardia minutissima consists of circum scribed reddish brown slightly scaling patches in the axillae and the

groins

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Histopathology For the demonstration of fungi in histologic sections the periodic acid Schiff stain of Hotchkiss McVanus is of very great value since it stains fungi deeply red while almost all other structures stain a very pale pink (kligman and Mescon). It should always be used when the presence of fungi is suspected. The reason for the strongly positive reaction of fungi to the periodic acid Schiff (PAS) stain is that the cellwalls of fungi are composed of mixtures of cellulose and chima which are rich in follysacharides (kligman Mescon and Delamater).

In histologic sections Jungi may present two structures hyphae (or mycela) and sporesy Hyphae which are nonreproductive appear as threadlike structures they may be septate or nonseptate. The chores which represent reproductive cells appear as round bodies.

TINEA OF THE GLABROUS SAIN. In tine of the glibious skin fungion of the crustine organism Malasse in furfur is always present in small numbers with the exception of tine versicolor in which the crustine organism. Malasse in furfur is always present in abundance. The fungi are found situated mainly in the upper two thirds of the stratum corneum. They may penetrate as far down as the upper layer of the stratum granulosum but they do not occur between thing cells. In fune of the glabrous skin except tines versicolor the fungi occur as short and long segmented and occusionally granular hyphae extending parallel to the surface xpores are usually not observed (Peck.) In tine Wersicolor, one sees not only hyphae but also spores.

In the absence of fungi no diagnostic picture is presented by fungus infections of the glabrous skin. In most instances depending on the degree of reaction of the skin to the presence of fungi one sees the histologic picture of either in icute 1 subscute of 2 chiquic demantits (see p. 68). Dermatophytosis of the hands and the feet

this with intra epidethiai 4000 Tinea Capitis and Tinea Barbae. In tinea capitis uncomment barbae fungs are present not only in the horny layer of the skin but also within and around the hair kligman, who recently has studied histologically the sequence of events in infections of human hair with Microsporum canis and Microsporum audouini found that

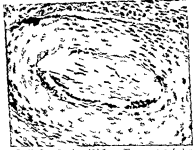


Fig. 100 Payus Periodic acid Schiff stain. The cross section of a hair follicle is shown. Il e lungus. Trichoplyton schoenleins is present largely as mycel a within and around the hair (X400)

after infection of the scalp surface. hyphae grow downward into the follicle on the hair's surface a short distance and then penetrate into the hair. Inside the hair they form branches and grow downward but not farther than the exact limit of the zone of keratin synthesis Thus as in the surface epidermis only fully keratinized structures are invaded Alter the invasion of the hair ectothrix spores are formed by segmentation of external branches of intrapilary hyphae Within the hair however the hyphae form no spores. As the infec tion subsides there is gradual reduction in the quantity of intra pilary hyphae and ectothrix spores

Aside from the presence of fungi the histologic picture in tinea capitis and tinea harbae may be merely one of subacute or chronic dermatitis Inserion Celsi and timea barbae however folliculitis the toes and an erythematous, scaling and vesicular eruption on the feet and the hands, especially the soles and the palms

Onychomycosis, caused usually by Trichophyton rubrum, shows

disintegration of the nul substance

Facus, rare in this country, is caused by Frichophyton (Achonon) schoonlein. Most commonly it affects the scalp, where it produces inflammation with formation of perifollicular crusts, called scittled Destruction of the hair ensues. Healing takes place with scarring

Tinea versicolor, caused by Malassezia furfur, most commonly affects the upper trunk, where one finds areas of brownish discolor tion. The surface of the discolored areas shows fine branny scaling

Erythrasma, caused by Nocardia minutissima, consists of circum scribed, reddish brown, slightly scaling patches in the axillae and the groins

Histopathology. For the demonstration of fungi in histologic sections, the periodic acid Schiff stain of Hotchkiss McManus is of very great value since it stains fungi deeply red while almost all other structures stain a very pale pink (Kligman and Mescon). It should always be used when the presence of fungi is suspected. The reison for the strongly positive reaction of fungi to the periodic acid-Schiff (PAS) stain is that the cellikulls of fungi are composed of initures of cellulose and chum which are rich in polysiccharides (Kligman, Mescon and DeLamater)

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Tiken of the Grandous Sain. In times of the glabrous skin, fungiare present in small numbers with the exception of times versicolor in which the causitive organism, Malassezia furfur, is always present in abundance. The fungi are found situated mainly in the upper two thirds of the stratum corneum. They may penetrate as far down as the upper layer of the stratum granulosum, but they do not occur between withing cells. In finite of the glabrous skin, except times versicolor, the fungi occur as short and long segmented and occasionally granular hyphie extending parallel to the surface exports are usually not observed (Peck.) In times Mersicolor, one sees not only hyphic but also spores.

In the absence of fungi, no diagnostic picture is presented by fungus infections of the glabrous skin. In most instances, depending on the degree of reaction of the skin to the presence of fungi, one sees the histologic picture of either an acute a subjective of a chronic degracius (see p. 68). Dermatophytosis of the hands and the feet

TABLE 4—Histologic Appearance of the Tissue and of the Yungi in Fungus Diseases

PLAGUS MISEASES				
Disease	Histologic Appearance of Tissue	AVERAGE Size of Fungus (Michons)	Appearance of Fungus by Tissue	
Monil asis	When invasive ponspecific granulation tissue with	4	Hyphae and budding as well as nonbudding yease	
Blastomycous	abscess formation Epithelial hyperplasia tu berculo-digranulation tis- sue with formation of	10	cells (spores) Thick walled spores in grant cells and tissue Budding forms	
Torulosis	Small abreesses Chronic inflammators infil trate with extensive cale	7	Spores with wide gelatin	
Chromobiastomytous		30	Thick walled, dark brown sporce, often in clusters Some cells possess tross	
Coccid aidom/cosis	Granulomatous nodules hise blastomy costy except that <u>casearion</u> may oc cur Subcutaneous ab- scenses central casearion autrounded by tuber		walle Thick walled spores with grandar syntplasm. The larger spores contain endospores.	
Actinomy costs	culoid granulation tissue onchecific granulation tis- sue with abocess forma non	150	Large irregularly lobu lated granules with branching filaments and club formation at the	
πρετοιπείου ισ	Primary lesion transpectific granulation tissue. Sub- cutaneous nodules thre zones chronic suppura- tive tuberculoid and syphiloid zone.		persphere Usually, no fungs are seen Occasionally, asteroid forms of spores are present	
H topla moss	Change granuluma way	3	*umerous spores sur rounded by a clear halo he in the cytoplasm of large histocetes	

ind foreign body grant cells. The infiltrate may extend into the subcutis. As a rule. Candida albicans is present only in the stratum corneum and not in the dermis (Hauser and Rothman).

In the rare instances of abscess formation and ulceration die we dudded additions hiphae as well as spores have been found in the determit expectably in the abscesses and their vector in. The presentation of micross in diameter, here a distinct current way.

and perifolliculitis (see page 168) frequently are seen At first, an and peritoniculitis (see page 168) frequently are seen At first, an intrafollicular abscess forms. The wall of the follicle may then rupture, whereafter the perifollicular tissue becomes the sent of an abscess. After discharge of the pus, the abscess cavity begins to fill in with granulation tissue. At that stage, epithelioid cells and foreign body guant cells are frequently present, digesting remnants of the follicular epithelium (McCarthy). It may be pointed out that the histologic pictures of tinea barbae and folliculitis barbae (sycosis sulgaris) are essentially the same (see page 169)

FAVUS The fungus is present in the horny layer of the skin, within and around the hairs (Fig. 105), and in the scutula. In some in stances, the fungus invades the stratum malpighii and even the dermis (Ormsby and Montgomery) The scutuli are composed it their periphery of well preserved hypline and spores while in the center one finds granular debris and degenerated spoies. The setuith rest upon an atrophic epidermis. The hair follocles underlying the scutula are destroyed. The dermis shows a mild to moderately severe inflammatory reaction

MONILIASIS

Candida albicans, the cruse of moniliasis, not only is a cutineous pathogen but also may affect mucosal surfaces, such as the ord mucosa and the vagina. In rare instances. Candida albicans may in fect the lungs or the meninges and cause death

Clinically, cutaneous moniliasis usually is a superficial infection characterized by fairly sharply demarcated areas of erythems with or without pustulation. The intertriginous areas are predominantly affected Paronychia is common In rare instances granulomatous and hyperkeratotic lesions form on the face and the sculp of children so called monthal granulous, and, in adults abscess formation and ulceration may occur

The association of cutaneous and mucosal moniliasis with hypo adrenalism (Talbot Butler and MacLachlan) and with hypoparathy roidism (Sutphin Albright and McCune) is known to occur Since this association is often familial, it is likely that both the endocrine deficiency and the susceptibility to monifol infection in these cases are due to some defect in the germ plasm

Histopathology. Superficial cutrineous monthrsis presents a histo logic picture like that found in subjectie or chronic dermitiis. The logic picture like that found in subscute or chronic definition. The fungus elements occur only in the stratum corneum. They consist of hybrid and spores, some of the latter in a budding stage. Monthal granuloma shows hyperkeratosis accuthosis and, in the dermis, a dense infiltrate of lymphocytes, plasma cells, neutrophils.

TABLE 4 —Histologic Appearance of the Tissue and of the I unglish

L'AGGS DISEASES				
DISEASE	Histologic Appearance of Tissue	AVERAGE SIZE OF FLAGUS (NICRONS)	APPEARANCE OF FUNGUS IN TISSUE	
Mon hasis	When invasive nonspecific granulation tissue with	4	Hyphae and budding as well as nonbudding yeast cells (spores)	
Blastomycosis	abscess formation Epithelial hyperplasia, tu berculo digranulation tis- sue with formation of	10	Thick walled spores in grant cells and tissue Budding forms	
Torulosis	small abscesses Chronic inflammators infil trate with extensive case ation	7	Spores with wide gelaun tous expende	
Chromoblastom\co-is		10	Thick walled, dark brown spores, often in clusters Some cells possess eross	
Coct d to domn costs	Granulomatous nodules hie blastomycoss except that <u>caseation</u> may oc cur Subcutaneous ab scesses central caseation surrounded by tuber	40	walls Thek walled spores with granular cytoplasm. The larger spores contain endospores	
Actinomycosis	culoid granulation tissue Nonspecific granulation tis- sue with abscess forma tion	150	Large, screpularly lobu lated granules with branching filaments and club formation at the periphery	
Sporatneho is	Primary leuon nonspecific granulation tissue. Sub- cutaneous nodules three zones chronic suppura- tive tuberculoid and syphiloid zone.		Usually, no fungs are seen Occasionally, asteroid forms of spores are present	
Histoplasmosis	Chronic granuloma wird foci of necrosis	3	Numerous spores sur rounded by a clear halo fie in the cytoplasm of large fistiocytes	

and foreign body giant cells. The infiltrate may extend into the subcuris. As a rule. Candida albicans is present only in the stratum corneum and not in the dermis (Hauser and Rothman).

In the rare instances of abscess formation and ulteration due to Candida albicans, hyphre as well as spores have been found in the derrmic especially in the abscesses and their vicinity. The spores measure about 4 microns in diameter, have a distinct capsule and

are Gram positive. They often lie in clusters and budding forms in occasionally seen. The hyphae are slender rods and have a beaded of segmented appearance (Rockwood and Greenwood, Danbolt)

NORTH AMERICAN BLASTOMYCOSIS (GILCHRISTS DISEASE)

North American blastomycosis, caused by Blastomyces dermatitudis is characterized by granulomatous and suppurative lesions which my occur in any organ of the body but are found most commonly in the <u>skin</u>, the <u>lungs</u> and the <u>bones</u>. Two clinical forms can be differentiated, a primary cutaneous and a systemic form. In the beings primary cutaneous form, one observes either one or a few-rather large lesious. In the datal systemic form, the cutaneous lesions are numerous and usually small in size.

The cutaneous lessons consist in both forms of <u>vertucous plujues</u> showing an active border beset with a large number of minute ab scesses. In the primary cutaneous form, the older lessons show central healing with atrophy. In the systemic form, the cutaneous lessons my undergo ulceration and, in addition, subcutaneous abscesses my occur.

Histopathology. Histologic evanination shows annihosis, papil lomatosis and considerable downward proliferation of the epidermis. The downward proliferation may be marked enough to present the picture of pseudo-epitheliomatous hyperplasia. Intra-epidermial absensesses are often present. Occasionally, one finds Langhans grant cells completely enclosed by the proliferated epidermis (Fig. 106).

The dermis is infiltrated by a polymorphous granulation tissue Polymorphonuclear leukocytes usually are present in large numbers and form small abscesses in the dermis. I anglians grant cells are settlered throughout the dermis. They usually he alone and not within groups of epithelioid cells. Occasionally, one observes tuberculoid formations but never true tubercles.

The cells, on spores, of Blastomyces dermatitudes are Furly numer ous in listologic sections. They are often found lying free in the issue, priticularly in the abscesses. They occur, however, in their largest number within the giant cells. One or several spores may be within a guant cell (1 ig. 107). When in this location, the spores are easily spotted, even at low magnification, in sections stained with routine stains. Being unstrined, they resemble smill, round hole punched out of the cytoplasm of the grant cells. On high magnification, the spores are seen to have a thick wall, which gives them a double contoured appearance. They measure from 8 to 15 microis in diameter, on the average 10 microis. Blastomyces reproduce by





Fig. 107 North American blastomycosis. High magnification of Fig. ure 106. Three blastomyces cells. (B.C.) are shown lying in the cytoplaim of giant cells. (X400).

budding, and occasionally <u>budding for</u>ms are seen in sections. Like in most fungus infections, many more spores are seen in sections stained with the <u>Hotchkiss McManus stain</u> than in routinely stained sections. (see page 218)

The lustologic appearance of the isserial lesions is unilogous to that of the cutaneous lesions. The number of neutrophils is often great and numerous abscesses tend to be present (Litiman, Wicker

and Warren)

Differential Diagnosis. Tuberculosis verrucosa cutis, torulosis and chromoblastomycosis must be considered in the differential diagnosis Tuberculosis vertucosa cutis shows no epores in the tissue. In addi tion, the number of neutrophils is much smaller and one usually finds true tubercles and areas of caseation necrosis. The distinctive features of torulosis and chromoblastomycosis are discussed under these re spective headings (see below)

FORULOSIS (CRYPTOCOCCOSIS, LUROPFAN BLASTO MYCOSIS OF BUSSE AND BUSCHKI)

Torulosis, though very rare, occurs throughout the world. The dis case, caused by Cryptococcus neoformans (Forula Justolytica), usually is chronic and systemic and ends in death. The main and the menin ges are nearly always affected, resulting in meningo encephalitis with presence of the organisms in the spinal fluid. The lungs also me commonly involved Cutaneous lesions are present in about 10 per cent of the cases, and, in rare instances, lesions of the oral mucosa occur (Cawley) The cutaneous lesions may be the first clinical sign of the disease (Linell, Magnusson and Norden), and in some instances the disease may remain limited to the skin (Gandy) In that case, the course of the disease is benign and self limited

The occurrence of torulosis has been reported either as a silent or as an active infection in patients with various types of lymphoma such as Hodgkin's disease (Geudel, Ende and Norman) and various types of leukemra (Cawley) \ similar association with lymphoma

Occurs also in histoplasmosis (see pige 231)

The cutaneous lesions are variable and may consist of pap ules, pustules, nodules, infiltrated plaques, ulcers or subcutaneous

abscesses

Histopathology. Histologic examination of the skin shows, in most instances, a chronic inflammatory infiltrate with or without giant cells Occasionally, the inflammatory reaction is mild and extensive caseation is present in the dermis

The causative fungus usually is present in abundance, either within

giant cells or lying free in the tissue. It consists of a spherical spore mension of the multiplies like. Blastomyces dermaintalis by building. It usually is surrounded by a wild gelationus capsule which does not stain with hemitoxylin and mere it stains ted with the persoduc acid Schiff reaction, and mere "Clanell Ving." (Linell Ving.

sule is absent

(Wile) The langus then greiny reschings 4 , *s dermainths and earth can be confused with it especially since both may octur in gain cells. On cultures however in contrast with Blastomyces Torolds forms no mycelas (Benham)

CHROMOBLASTOMYCOSIS

This discuse is hanted to the skin and is benign. It is crused by three closely related fungi which appear alike in the tissue. Hormo



Fig. 103 Chromoblastomycous Petrodic acid Schill stain. Epithelioid cills and Langhains giant cells from the 1-all of an abscess. Two long et aum of lungus cells are located in the right upper corner. In addition three lungus cells he in the right center. (x400)

dendrum pedrosos Phialophora verrucosa and Hormodendrum com pactum (French and Russell)

The curvaeous lesions usually limited to a single area consist of densely aggregated nodules and plaques with a hard verrucous surface

Histopathology. The histologic appearance of the skin is very much like that of blastomycosis. The pidermis shows considerable hyper plasia although rarely to the extent that is usual in blastomycosis. The vermis shows an intensite infilitation, with a polymorphous granulation tissue containing many Langhans guant cells and small abscesses composed of polymorphonuclear leukocytes. Tuberculoud formations may be present, but true tubercles and case tion are absent, just as in blastomycosis.

The causitive organisms are found in Lingham giant cells is well as free in the tissue and are especially numerous in the abscesses. They appear as sclerotic, dark brown, thick-walled, usually spherical spores varying in size from 8 to 15 microns. They he either singly or in chains or clusters (Fig. 108). Reproduction is by intracellular will formation and splitting not by budding. In some of the organisms cross walls can be seen.

Differential Diagnosis. Chromoblastomycosis cannot be differentiated from blastomycosis except through the different appearance of the fungi-Just as blastomycosis chromoblastomycosis differs from tuberculosis verrucosa cutis by the presence of a fungus in the tissue, and the absence of true tubercles and of criseation

COCCIDIOIDOMYCOSIS (SAN JOAQUIN VALLEY FEVFR)

Coccidioidomycosis is caused by Coccidioides immitis. It is endemic in the southwestern United States especially in the Sin Jorquin valley of California, and in northern Mexico. Three forms are recog nized, primary, intermediate and progressive (Duemling)

Timary coccidioidomycosis represents an acute respiratory infection. Development of erythema nodosum is not uncommon

Progressive coccidioidomy.cosis (coccidioidal granulomi) tolions the primary form in a small percentage of cases after a varying length of time. It has a very high mortality. Many organs especially the meninges, the lungs the bones and the lymph nodes may be in volved. Cutaneous lesions are common. They consist either of ver rucous and granulomytous nodules or of subcutaneous cold abscesses which may break through the skin.

Histopathology. The nodose lesions of armary coccidioidomycosis have the same histologic appearance as idiopathic erythema nodosum (Winer)

The verrucous and the granulomatous lesions of coccidioidal granuloma resemble blastomycosis in their histologic aspects. How

ever there is less tendency to abveess formation, and caseation necrosis may octur (Moose). The causative organisms are found free in the ussue as well as in Langhans-giant-cells. As a rule they are present in lunge number.



Fig. 109 Coccidic larger than those o size Their cytoplas

The subcutaneous abscesses of coccidioidal granuloma resemble scrofuloderma in their last logic and an analysis of the second of the subcutaneous abscesses of coccidioidal granuloma resemble scrofuloderma in their last logic and the subcutaneous abscesses of coccidioidal granuloma resemble scrofuloderma in their last logic and the subcutaneous abscesses of coccidioidal granuloma resemble scrofuloderma in their last logic and the subcutaneous abscesses of coccidioidal granuloma resemble scrofuloderma in the subcutaneous abscesses of coccidioidal granuloma resemble scrofuloderma in the subcutaneous abscesses of coccidioidal granuloma resemble scrofuloderma in the subcutaneous abscesses of coccidioidal granuloma resemble scrofuloderma in the subcutaneous abscesses and the subcutaneous abscesses are subcutaneous abscesses and the subcutaneous abscesses are subcutaneous abscesses and the subcutaneous abscesses are subcutaneous abscesses are subcutaneous abscesses and the subcutaneous abscesses are subcutaneous abscesses are subcutaneous abscesses and the subcutaneous abscesses are subcutaneous are subcutaneous absc



Fig. 110 Coccidioidomycosis. A large Coccidioides spore lies within a grant cell. The Coccidioides spore contains numerous endospores (X400).

thehord cells and some grant cells. Numerous spores are present extracellularly as well as intracellularly in grant cells.

The spores of C.

80 microns

10 to

Coccidioides is much larger than Blastomyces, 1 orula or Phialophora The spores are spherical and thick walled and hive a granular cytoplasm. Multiplication is not by budding but by formation of endo. Spores which may be seen lying inside the larger spores (Lig 110). The endospores are released into the tissue by rupture of the wall of the spore. Indospores may measure up. to 5 microus in diameter.

Differential Diagnosis. A diagnosis of coccidioidomycosis can be made only in the presence of the fungus

ACTINOMYCOSIS

Actinomycosis is caused either by Actinomyces boyis, which is anaerobic or nucro aerophilic, or by several species of Nocardia (eg N asteroides, N madurae), which are aerobic

Actinomycosis frequently affects the skin Involvement may be primary, as in Madura foot (injectoma) More often, however, the infection reaches the skin from an internal focus. The most common form is vervicofacril actinomycosis, in which case the organism reaches the skin from the mouth, next in frequency are actinomy costs of the thoracie skin secondary to involvement of a lung and actinomycosis of the addominal skin secondary to involvement of the occurrence of the appendix.

The involved in is dark red in color, possesses a "wooden tipe of hardness and shows numerous sinuses discharging a seroanguine ous or purulent fluid containing suffur yellow granules which consist

of masses of fungi

Histopathology. Histologic examination shows extensive granulation tissue containing large abscesses. The fungus granules are found within the abscesses. The canulation tissue is nonspecific in its appearance. In the early phase of the disease it is composed of neutro phils, cosmophils, lymphocytes plasma cells shistocytes and sfibro blasts. In the healing phase fibroblasts predominate. Thus, the dreg nosis can be established only by finding the Actinomyces granules in the abscesses. When selecting an area for biopsy, an area containing purulent material should be chosen.

They ungus granules are large and may measure several hundred microns in diameter, large enough to be visible macroscopically 18 the so called sulfur granules. In histologic sections, they appear base (Fig. 111). They are homogeneous granching filaments at the periphery

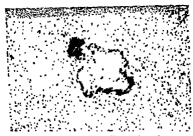
The ends of the filaments requently are surrounded by a gelatinous sheath, giving the ends a club shaped appearance. The filaments are much better seen in sections stained by Grain's method, which stains them Gram positive, than in sections stained with routine stains. The

cells immediately around the granules are usually polymorphonuclear lenkocytes, but foreign body grant cells occasionally are seen in

contact with the granules

Sunr .- "

In general, Nocardia produces more Inecrosis and less granulation tissue and less fibrosis than Actinomyces The granules of Nocardia are smaller and much less numerous than those of Actinomyces so that multiple sections are necessary to find them (Weed and Baggen



accommended A large sulfur granule is shown in the center of the held. The granule appears homogeneous in the center Clubs are present at the periphery (X100)

stoss) Nevertheless, these differences are not sufficiently distinct and cultural studies are necessary for a differentiation of Nocardia from Actinomyces

SPOROTRICHOSIS

mkn, affects only the benign disease Most

type there is a primary lesson (sporotrichout chancre) which is commonly an ulcer but may be papillomatous Secondarily, multiple sub cutaneous nodules appear along the kanphance des -area The nodules ---

Much ... unit are widely scattered lesions consisting seminate the localized type, form

Coccidioides is much larger than Blastomyces, I orula or Phialophora The spores are spherical and thick walled and have a granular cyto-plasm Multiplication is not by budding but by formation of endo, spores which may be seen lying inside the larger spores (Lig 110) of the endopores are released into the tissue by rupture of the will of the spore Endospores may incasure up to 5 microus in drimeter Differential Diagnosis A diagnosis of coccidioidomycosis can be

made only in the presence of the fungus

ACTINOMYCOSIS

Actinomycosis is caused either by Actinomyces bows which is amerobic or nucro aerophilic or by several species of Wocardia (e.g.

N asteroides N madurae) which me nerobic

Actionary of managery which the service.

Actionary of the facts the skin Involvement my be primary as in Vidura foot (mycetoma) More often however the infection reaches the skin from an internal focus. The most common form is very cofresil retinomy costs in which case the organism reaches the slan from the mouth next in frequency are actinomy costs of the Morrace slan secondary to involvement of a lung and actinomy costs of the Morrace slan secondary to involvement of the cecum or the appendix

The involved kin is drik ted in color possesses a wooden type of hardness and shows numerous sinuses discharging a serosanguine ous of purulent fluid containing sulfur vellow granules which consist

of masses of fungi

Histopathology Histologic examination shows extensive granula tion tissue containing Juge phocesses. The tringles granules are found within the abscesses. The trinulation tissue is nonspecific in its ap pearance. In the early phase of the disease it is composed of neutrophils eosinophils elymphocytes plasma cells ehistiocytes and fibro blasts In the healing phase fibroblasts predominate Thus the day nosis can be established only by finding the Actinomyces granules in the abscesses When selecting an area for biopsy an area containing purulent, material should be chosen

The fungus granules are large and may measure several hundred microns in drumeter large enough to be visible macroscopically as the so called sulfur granules In histologic sections they appear base
111) They are homogeneous

un, filaments at the periphery

The ends of the filaments requests to the periphers sheath giving the ends a club shaped appearance. The filaments are much better seen in sections stained by Gram's method which stains them Gram positive, than in sections strained with routine stains. The

thick walled round spores are found free in the tissue visible even with hematoxylin cosin strining. Some of them have radiating as teroid clongations (Fig. 152) (Moore and Ackerman. Pinkus and Grekin's

Differential Diagnosis Without the demonstration of the Jungus a diagnosis of spototrichosis cannot be made but can only be sus pecied. The subcutaneous nodules of tularemia may have the same histologic appearance Explient industrium differs from sporour chosts by its vascular changes and the presence of caseatton which is absent in sporottichosis

HISTOPLASMOSIS

Histoplasmosis caused by the lungus Histoplasma capsulatum in most instances is a fatal systemic disease. Occasionally the disease is limited to the sem or the mucous membranes and benign in its course.

infinite to the section in the management of the section in the section of histoplasmons the clinical picture is sarable. The lymph nodes are often enlyinged markedly. Pullmonary and adversy modification they be a prominent feature. Cutaneous or mucomembranous lessons occur in about half the princips with the systemic form of infection (Miller Keddie Johnstone and

Several authors have reported the simultaneous occurrence of I stonl mn nd

sease (Fnde and Curus)

us , to the lym

thoma especially in sien of the fact that also torulous has been chierced in association with lymphoma (see page 224).

The skin lesions of histoplasmosis in both the systemic and the

cutaneous form may be nodulat granulomatous or olcerative

Histopathology Histologic examination reveals a chronic non specific granulomatous infiltrate with foci of necrosis Throughout this infiltrate but especially in the ricinity of the necrotic foci one observes large histocytes (macrophages) engorged with numerous organisms

Histof lasma capsulatum appears in sections and with hema surrounded by

organism meas L to a uncrons in diameter. On training with the periodic acid Schill reaction however Histoplasma capsulatum shows no cap sule but instead the body appears larger and lined by a definite red stained cell hall. Therefore, it appears that the so-called halo or Histopathology. The primary lesion of sporotrichosis shows a non specific granulation tissue containing, among other cells, mun plasma cells, epithelioid cells and some Langhaus giant cells. For of suppuration are present. If the primary lesion is pipillomatous, one finds, in addition, marked epithelial hyperplasia and intra epidermal abscesses (Moore and Ackerman)

The subcutaneous nodules show a more characteristic appearance than the primary lesion because of the arrangement of the infiltrate

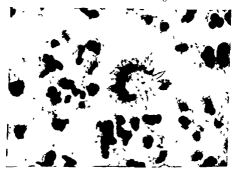


Fig. 112 Sporotrichosis \ large spore with radiating asteroid clongs tions is shown (×1850) (Hermann Pinkus M D)

in three zones—the chroms suppurative zone, the tuberculoid zone and the syphiloid zone. The zone in the center of the nodule, the chrome suppurative zone is composed chiefly of neutrophils with a few histocytes and lymphocytes. Small abscesses composed entirely of neutrophils are present within this zone. The middle or tuberculoid zone is characterized by numerous epithelioid cells and a large number of grant cells of the Linghaus type. The grant cells may lie in groups. The peripheral or syphiloid zone consists of a righty cellular infiltrate of plasmicells, lymphocytes and fibroblasts. The zonal arrangement is not always distinct. In older nodules one merely sees a nonspecific granuloma.

The causative organisms are present in the tissue in small numbers as round to fusiform bodies which do not stain with hematoxylin and cosin but become visible when the periodic acid Schiff reagent is employed (Kligman and Baldridge). In rate instances, a few large,

233

Tomiosis

Benham R W. The fungi of blastomycosis and coccidioidal granuloma. Arch Dermat & Siph 30 385 1934 Cawley E P Grekin R H and Curtis \ C Torul sis J Invest. Dermat

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special reference to apparent association with Hodgkin's disease Am J Med 9 343 1950 Linell F. Magnusson B. and Norden A. Cryptococcosis. Acta dermat venereol

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cosa) 1 Cutan Dis 33 810 1915

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Kligman \ M and Baldridge G D Morpholom of Sparotrichum schencku

A comparison of to (Good reason) rmation in tissue

and kile R L. Generalized subcutaneous gummatous ulcerating sporotrichosis, Arch Dermat & Syph 31 672 1935

Pinkus II and Grekin J > Sporotrichous with asseroid tissue forms Arch Dermat & Syph 61 813 1950

<u>censule</u> is an artefact produced by plasmolysis which causes the eyo plasm to shrink away from the cell will (Migman and Baldridge) *Histoplasma capsulatum* is a fungus according to its cultural that acteristics and because it multiplies by building

Differential Diagnosis The general appearance of the granulomatous infiltrate with its parasitized histocytes is much like that of rhunoselcroma, granuloma inguinale and cutaneous leishimanisis (Lor their differential diagnosis see Table 5 page 237)

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epidermis grafted on the chorio all intois of chick embryos with the fluid obtaiged from herpes zoster vesicles

It is probable that the viruses of varicella and herpes zoster are identical. There exists strong clinical evidence (Bruungaard) and experimental evidence (Braun) for this contention Turthermore the two viruses have an identical appearance under the election microscope (Baldrudge Blank and Rake). Wise and Subberger beheve that herpes zoster represents an unmune or allergic form of varicella which occurs in patients who had previous clinical or subclinical varicella.

varietla
Clinical Appearance variola shows a generalized eruption which
at first consists of papules. After two or three days, the papules are
transformed into vesicles which are characteristically imbificated.
After three more days, the vesicles change into possibles. In severe
cases in addition there may be purpoure lesions. It is characteristic
of variola that all leagues are at the same stage of development.

The etuption of varicella is also generalized The lesions begin as small papules which soon develop into <u>resides</u>. The vesicles crust over as a rule without changing intopustules. Lesions occur in successive crops so that one observes lesions in different stages of development.

£

hota uttu eruption there is a generalized vesicular eruption indistinguishable from that of varicella. Herpes zoster (especially the generalized form) ys apt to occur in lymphoma (see page 491)

Herpesymplex shows one or several groups of vesicles on a mildly inflammatory base. The eruption may occur anywhere on the skin but is found most commonly about the face, and the genitalia

Histopathology The characteristic histologic lesion in the four diseases is an intra epidermal vesicle produced by profound degeneration of epidermal cells Because of the presence of these degenerative changes the vesicle differs histologically from those seen in other

of Bullae page 66) and et it is important that as ind be selected for biopsy

invasion of inflammatory epidermal cells occurs in two forms ballooning degeneration and reticular degeneration.

Ballooning day -

cosinophilic cytoplasm (Figs. 114, 115). They may have no nucleus, or may have one of many. Because of the fact that the balloon cells lose their/intercellular budges, acantholysis occurs and the cells be come separated from one another. Unilocular vesicles result. The process of ballooning degeneration occurs mainly at the base of virus vesicles leading to a dissolution of the lower epidermis, so thit, ulti



Fig. 114 Herpes zoster. I on mignification. There is marked billioning degeneration of the cells at the floor of the visitle. The cells of a fur follieft, shown it the left, likewise show billioning degeneration. Reticular degeneration observed at the top of the vesicle, is only slight since this is an early lesion, no inflammatory reaction is present (×100).

mately, the originally intra-epidermal vesicles become subepidermal in many places. Ballooning degeneration affects also the epithelial cells of hair follicles and bebaceous glands.

Keticular

the cell wall bursts. By coalescence of neighboring, similarly accells, a multilocular vesicle results, the septa of which are formed by the resistant cellular walls (Fig. 116). Reticular degeneration occurs mainly at the top and the periphery of virus vesicles. In older vesicles, the resistant cellular walls disappear and the multilocular vesicle then becomes unilocular. It may be pointed out that reticular degeneration is not/specific for virus vesicles since it also occurs in the vesicles of dermatius (see page 68).

numerous polymorphonuclear leukocytes as well as some macrophages (Sheldon and Heyman) These central abscesses tend to have a triangular or quadrangular shipe with elongited corners, giving them a <u>stellate appearance</u> (Fig 124) The epithelioid cells surround ing the abscesses are arranged in palisade formation. As the abscesses gradually enlarge, they coalesce and lose their stellate appearance



to 121 symphogramitoms renereum. In the lymph node, a stellate abscess is present. It is surrounded by epithelioid cells in palisade formation (×100).

MILKERS' NODULES

Millers nodules are acquired from constintened with paravac cuma intural compox? There are, usually on the Ingers, one to three and occasionally more nodules, I to 2 cm in diameter, which are blush red, semiglobular and usually <u>printers</u> Spontaneous healing occurs in I to 2 months

Histopathology. The Spiderims shows acustings and parakera tons, A dense wonspecific chronic inflammatory infiltrate is present in the derims. The capillaries are increased in number, are dilated and show swelling of their endothelial cells (Nomland and McKee). While most authors have found no inclusion bodies, Kattenellen.

bogen observed them in some but not in all of his cases. They were

located in the cytoplasm of vacuolated epidermal cells and often were Feulgen positive. Their appearance was similar to that of the inclusion bodies observed in variola and vaccinia.

Study of smerrs of tissue obtained from milkers nodules under the electron microscope has shown the presence of typical elementary bodies of the paravaccious type (Nascimann and Deubner)

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Study of smears of tissue obtained from milkers nodules under the electron microscope has shown the presence of typical elementary bodies of the paravaccinia type (Nasemann and Deubner)

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1940

16

Metabolic Diseases

LIPOIDOSES

The term lipoidoses has been applied to a group of diseases in which the lesions, due to a local or generalized disturbance of the lipid metabolism, contain lipid substances.

No generally accepted classification of the lipoidoses exists. The following classification is based largely on the classifications published by Thannhauser and by Montgomery and Osterberg

- t Lipoidoses with Increased Blood Lipids
 - Primary hypercholesteremic xanthomatosis
 - 2 Biliary xanthomatosis
 - 3 Idiopathic hyperlipemia
- 4 Secondary hyperlipemia II Lapoidoses with Normal Blood Lapids
 - 5 Land reticula endotheliosis (Hand Schuller Christian disease)
 - (a Fulminating type Letterer Sine disease Jh Remi her on 11 -161
 - - 8 Lipoid proteinosis 9 Extracellular cholesterosis
- III Localized Lipoidoses 10 \amhelasma palpebrarum
 - 11 Necrobiosis lipoidica

LIPOIDOSES WITH INCREASED BLOOD LIPIDS

1 Primary Hypercholesteremic Vanthomatosis Primay

managaeuna tat Therefore the serum is

Entaneous lessons consist of flat or slightly raised xanthelasmata on the eyelids and tuberous xanthomata, especially on the elbons and the knees. In addition, there are tendon canthomata, especially on the Achilles sendons, the patellar tendons, and the extensor ten

Diseases Caused by Viruses

1952

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in xanthoma cells to be anisotropic (doubly refractile) in contrast with the fat in the sebaccous cells and the subcutaneous fat cells, which is isotropic (not doubly refractile) (See page 31)

Yanthoma cells form from perithelial cells, which are histocytes
They have usually one nucleus but may have two and even many



Fig. 126. Aanthoma tuberorum early lesson. Scarlet red stain for fat. The xanthoma rells are filled with lipid material (×100).

nules in multimiciated xanthoma cells the nuclei either are irregularly distributed as in foreign body grint cells, or he near the center of the cell grouped around a small island of nonfoamy cytoplasm and surrounded by foamy cytoplasm. This bears

particularly polymorphonuclear leukocytes symphocytes and instacettes in belif-developed lessons the inflirate is composed almost entirely of xanthoma cells (Fig. 125). Fat stains in this stage show

Metabolic Diseases

258

dons of the hands Atherosclerotic cardiovascular disease is common and may lead to death due to coronary occlusion in early life (Bloom Kaufman and Stevens)

Histopathology. The histologic appearance of the cut; neous and the tendinous lesions is characterized by the presence of anthoma or foam cells. Xanthoma cells are phagocytic cells filled with lipid drop



Fig 125 Nanthoma tuberosum, early Iesion Numerous xauthoma cells (foam cells) are present There is only little fibrosis (×200)

lets In routine sections, the lipid droplets have been dissolved and extracted in the process of fixation and embedding so that the cells have a reticulated or framp appearance (Fig. 125). However, the lipid droplets can be seen when formalin fixed frozen sections are stained with far stains, such as scriterized (Fig. 126). With scarlet red, the lipid substance in the nanthomy cell stains a brownish red, in contrast to the lipid substance in schiecous glands and the subcutaneous fat, which stains a bright-orange red. This is due to the fact that the lipid substance in nanthoma cells is predominantly choles terol and phospholipids, whereas the lipid substance in the sebactous cells and the subcutaneous fat cells is predominantly neutral fat Polariscopic examination of frozen sections reveals the lipid droplets

xanthomatous lesions often by many years. The blood serum is clear but due to its high content of bilirubin it is intensely green

The biliary cirrhosis which causes the xanthomatous lesions is produced as a rule either by cholangiolitis or by extrahepatic ob struction (MacMahon) but occasionally in children by congenital hypoplasia of bile ducts (MacNahon and Thannhauser)

Histopathology The cutaneous lesions show the same histologic picture as is found in primary xanthomatosis

3 Idiobathic Hyperlipemia

In inopathic hyperhipemia which usually is not familial the blood serum shows an elevation not only of the cholesterol and phospholipids as in oriman hyperchalming.

also of the neutral fat Therefore the blood serum is mulky

The cutaneous lesions are of two types Muberous xanthomata which are found especially on the elbows and the knees and papular xanthomata which may be diffusely distributed but are most preva lent on the buttocks. The papular xanthomata have a tendency to come and go and therefore are often referred to as eruptive xanthomata Nanthelasmata of the eyelids are absent but Tendon xantho mata may occur (Lever Smith and Hurley)

Asceral manifestations occasionally are present. They include hepitosplenomegaly attacks of abdominal pain due to secondary pancreatitis and coronary heart disease which however occurs less frequently than in primary hypercholesteremic santhomatosis. The association of idiopathic hyperlipemia with hepatosplenomegaly oc

casionally is referred to as Burger Grutz disease

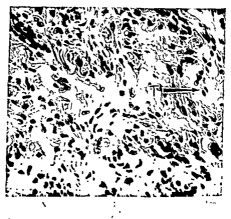
Histopathology The histologic appearance of the xanthomata in idiopathic hyperlipemia is the same as that of the vanthomata in primary hypercholesterenat xanthomatosis. The papular vanthomata which usually are of vecent origin often show like most young canthomata an admixture of inflammatory cells (see page 239)

Only lew reports on the histology of the Visceral lesions are avail able in one patient with hepatosplenomegaly conspicuous fat in filtration of the liver cells was observed on biopsy (Movitt Gerstl Sherwood and Epstein) The presence of fancreatitis in patients with attacks of abdominal pain has been confirmed repeatedly by explora tory operation (Klatskin and Gordon)

4 Secondary Hyperlipemia

Secondary hyperhipemia may occur secondary to severe diabetes (vanthoma diabeticorum) toonephrosis and to ellicogen storage disease (son Gierke's disease) The amounts of cholesterol phospho

that all fat is intracellular. In involuting lesions, fibroblasts appear (Fig. 127). Ultimately, fibrosis replaces the foam cells. Weidman and Schaffer express the belief that the foam cells themselves can become transformed into fibroblasts. In old fibrosing lesions cholesterol may be found not only in foam cells but also extracellularly.



center of the cell grouped around a small island of nonformy cyto blasm and surrounded by foam, cytoplasm (×400)

Differential Diagnosis. Differentiation between a fibrotic lesion of xanthoma tuberosium and histocytoma may be very difficult and even impossible. It may be necessary to tely upon-clinical and blood chemical data for the correct diagnosis (Montgomer) and Osterberg)

2 Biliary Nanthomatosis,

Biliary cirrhosis may cause high values for cholesterol and phos pholipids in the serum and lead to vanthelasmata on the eyelids and tuberous xamihomata on the skin that are indistinguishable from those of primary hypercholesteremic vanthomatosis. However, deep laundice is present and the jaundice precedes the appearance of

vanthomatous lesions often by many years. The blood serum is clear but due to its high content of bilirubin it is intensely green

The biliary citribosis which causes the xanthomatous lesions is produced as a rule either by cholangiolisis or by extrahepatic ob struction (MacMahon) but occasionally in children by congenital hypoplasia of bile ducts (VacVahon and Thannhauset)

Histopathology The cutaneous lesions show the same histologic picture as is found in primary xanthomatosis

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lipids and neutral fat in the serum are elevated and the serum is milky.

Papular eruptive vanthomata occur which usually are diffusely dis

tributed but have a predilection for the buttocks Occasionally, re

cently erupted papules have an inflammatory base

Histopathology. The pipular xanthomata of secondary hyper lipemia show the same histologic picture as other xanthomaty Since they usually are of recent origin, a rather marked inflammatory in filtrate may be present with large numbers of neutrophils In the wholuting stage, much of the cholesterol may be seen in phagocytes and extracellularly (Montgomery and Osterberg)

LIPOIDOSES WITH NORMAL BLOOD LIPIDS

5 Lipid Reticulo endotheliosis (Hand Schuller Christian Disease)

Letterer Siwe disease, Hand Schuller Christian disease and eosinophilic granuloma represent variations in degree, stages of develop ment and localization of the same basic disease process (Larber) For merly, Hand Schuller Christian disease was regarded as a primary disturbance of the lipid metabolism, as a normocholesteremic vin thomatosis (Thumhauser and Magendantz), but this concept has been abandoned (Mallory) and the presence of cholesterol in the lesions is now regarded as a secondary infiltration. The three diseases are at present believed to be histocytoses or reticulo endothelioses. If the reticulo endotheliosis occurs in Milancy, it is generalized and rapidly fatal (Letterer Siwe disease), death occurs before sufficient time has clapsed for the development of the losion into adipogrami loma In early childhood, the disease is Chronic (Hand Schuller Christian disease), and budization is as a rule pronounced In Inter childhood or in the idult the usual picture is that of eosinophilic granuloma which represents an abortive form of the disease Tran sitional cases between these three forms of reticulo endotheliosis are common The values for blood plasma hpids are normal in all three forms

LITTERER SIWE DISEASE (NONLIPID HISTIOCYTOSIS OF FOOT AND OLCOTT) This disorder usually occurs in infines and is almost in exitably fatal within a few months. It is virtuacterized by fever, anemir, enlargement of the liver and the spleen, lymphydenopythy and multiple defects of the bones. In most cases virtuacities lesions are present. They may consist of peticlinae pipules or pustules in some cases one observes numerous closely set, brownish pipules covered with scales or crusts. This type of cruption usually is extensive with a predilection to involve the scalp, the face and the trunk

The resemblance of the eruption to seborrheic dermatitis and

Daner's disease is often striking (Laymon and Sevenints)

Histopathology The cutaneous lesions show usually close to the epidermis and often invading into the epidermis (Fig. 128) accumulations of histocites (reticulum cells) intermingled with a few fem phocytes and varying numbers of cosmophils Extravasated blood cells frequently lie in and about the masses of hisnocytes. The his tiocytes appear as large cells with irregularly shaped vencular nuclei and abundant slightly eosinophilic cytoplasm (Fig. 129). In some areas these cells are distinctly outlined and even separated by edema but in other areas their cytoplasm is confluent Occasionally some of the cells have a form cytoplam and stam posture for far neth tat stants. The questions may become destroyed by pressure of the underlying cells (foot and Olcott. Abr and Denenholz. Line and Smith case I)

The Miceral lesions consist of proliferations of large pale reticulo endothelial cells which invide and replace the normal structure of spicen fiver bone marron lymph nodes and other organs (Sweitzer and Laymon)

HAND SCHLLLER CHRISTIAN DISEASE Diabetes insipidus exoph thalmos and multiple defects of the bones especially of the eranium represent the triad of typical Hand Schuller Christian disease. How ever any one or even all three of the cardinal symptoms may be absent and involvement may occur in entirely different organs. For example enlargement of the liver the spleen and the lymph nodes is common and dwarfism is observed occasionally. Hand Schuller Unistian discuse takes a chronic course usually extending over years and has a mortality of about 70 per cent

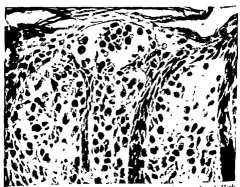
Cutaneous lesions are quite uncommon. If present, they are similar to those seen in Letterer Sine disease consisting of an extensive eruption of coalesting scale or crusted papules with a distinct clinical resemblance to Darier's disease Occasionally the term xanthoma dis seminitum is used for this eruption

Histopathology Early cutaneous lessons of Hand Schuller Chris tian disease show the same histologic picture as the cutaneous lesions of Letterer Sine dise use (Lane and Smith Laymon and Sevenants) Invitature lesions a large number of the histocytes show a foams cytoplasm and typical foam cells may be present (Thannhauser and Magendantz) In old lessons the number of foam cells again may be small and foreign body grant cells may be found As a rule the tendency to lipidization is less pronounced in the cutaneous lesions

than in the lesions of other organs



Lie 128 Hand Schuller Christian disease (cosinophilic granuloma) Low magnification. The upper portion of the dermis contains an infaltrate composed almost entirely of loosely aggregated distinctives. The infiltrate has invided the goldermis in many areas (x100).



ragnificatic
clet and abundant (×400)

loma) High shaped nu invaded the

Eosinophii ease of this gi

They occur moded The cutaneous lesions consist either of an extensive eruption of crusted papules as in Letterer Siwe and Hand Schuller Christian disease (Lever and Leeper) or of one or several erythemitous granulomatous plaques whigh may undergo ulceration (Curtis and Caules McCreary) The Mo types of lesions may be present simultaneously (Granuloma faciale formerly called eosinophilic granuloma of the face is an entirely different disease from eosinophilic granuloma and is not relayed to it. For its description see page 111.)

Histopathology The papules have the same histologic appearance as in Letterer Sine disease namely an infiltrate of loosely aggregated is and invading the epi.

The number offcosino

The frantiomatous plaques also show numerous loosely urranged large histocytes within an edematous strong Econophils are present in varying numbers. They usually he in packies rather than diffusely distributed through the lesion. In addition a few lymphocytes and plasma cells are present (Curtis and Cawley McCreary Lever and Leeper).

XLipid is absent in the cutaneous lesions but may be present in the

6 Atemann Pick Disease

Niemann Pick, disease is characterized by abnormal deposits of sphingamethr a diaminophosphatide in the reticulo-endothelial cells of many organs but not in the hard. The level of lipids in the bloods/serum is normal

The disease occurs as a rule in Jewish infants and is fatal. There are enlargement of the liver and the spleen cacheria and brownish discoloration of the skin

Histopathology On histologic examination the brownish discolor ation of the skin is found to be due to the presence of increased amounts of hielanin

7 Gaucher's Disease

In Gauci er's disease kerasin a cerebroside is deposited in the reticulo endothelial cells of many organs. The skin however is spared. The bloodylpids are normal

The disease tends to le familial, occurs predominantly in Jews may start at any age and takes a chronic course. There is hepatosple

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nomegaly and rarefaction with cortical thickening of the long bones. The skin shows brownish discoloration

Histopathology The skin shows increased amounts of melanin

8 Lipoid Proteinosis (Urbach II iethe)

Lipoid proteinosis is characterized by lipid infiltrations of the skin of the oral nucosa and of the largin. Although theyblood lipids may



Fig. 130. Lipoid proteinosis. The dermis is occupied by thick ways hydin like bundles which run perpendicular to the epidermis. In addition, the hydin like material surrounds all blood vessels as a thick mantle. (X200)

be normal (Wise and Rein Price LaRosa and Settle) there may be an increase in the total lipids (Hansen Ramos e Silva) or a relative increase in phospholipids (Montgomer) and Havens Wile and Snow) The disease is often familial

Climically, one observes nodular and vertucous lesions on the skin and on the mucous membranes of the mouth and the larynx. The nodules of the skin on regressing leave pitted scars, giving the skin a

I there is hoarse-

i is striking and irregular acan



Lipous proteinosis Scarlet red stain for fat. A large amount of lipid material is present. It occurs in the form of small droplets throughout the hyaline material, particularly around the blood vessels (x100)

thosis. The Mermis is considerably thickened. The upper half of the dermis is occupied by thick, navy bundles which are

homogeneous material is present around some of the vessels and sweat glands

On staining with scarlet red, a large amount of lipid material is visualized. It stains a rusty brown and occurs in the form of small droplets throughout the hyalin like material, but particularly around the blood vessels (Fig. 131). It is located extracellularly. On polari scopic examination, the lipid material proves to be isotropic not A doubly refractile Urbach and Wiethe interpreted the histologic find ings as indicating a merging of lipid and protein and, therefore, suggested the name lipoid proteinosis

No definite conclusion has so far been reached about the Ynture of the lipid substance in the dermis. In several cases, the chemical analysis of involved skin revealed a decided increase in the amount of lecithin, which is a monoaminophosphitide (Montgomery and Havens Ramos e Silva) In other cases, however, no such increase was found (Wile and Snow, Hansen, Price, LaRos) and Settle)

9 Extracellular Cholesterosis (Urbach)

This disorder is characterized by extracellular cholesterol infilira tions in the skin

Only three cases of this condition have been reported in the litera ture (Urbach, I'pstein and Lorenz Laymon Sobel and Pollock) All three cases showed reddish brown verrucous plaques and nodules involving especially the dorsa of the hands and the feet and the extensor surfaces of the legs Ulceration of some of the lesions occurred in Sobel's case. The amount of cholesterol in the blood/serum was normal in Laymon's case and subnormal in Urbach's and Sobel's cases

Histopathology. The involved skin shows a dense, nonspecific cel Inlar infiltrate composed mainly of histiocytes and lymphocytes, The blood vessels are dilated and their endothelium is swollen Voloam cells are present. In healing lesions, there is considerable librosis

Aat stains reveal numerous droplets of fat in extracellular position throughout the lesions but especially about the blood vessels. On polariscopic examination the fat droplets are doubly refractile

Chemical examination of tissue in Urbich's and Sobel's cases re vealed the content of cholesterol three to five times greater in the lesions than in the normal skin

The cause of the disease is unknown Sobel and Pollock impressed with the subnormal values for cholesterol in their as well as in Urbach's case, suggest that, for some unknown reason, the blood is unable to hold normal amounts of cholesterol which, therefore, is

deposited in the tissue. Being unable to metabolize the cholesterol the tissue responds not with foam cell formation but instead with a severe inflammatory reaction as if dealing with a foreign body of irritating or toxic nature.

LOCALIZED LIPOIDOSES

10 \anthelasma Palpebrarum

This disorder is characterized by the presence of soft yellowish plaques on the cyclids caused by the deposition of cholesterol Although xanthelasma pilpebrarum is common in primity hyper chofesteremic vanthomatous (see page 237) it frequently occurs in individuals with little or no elevation of the serum cholesterol (Epstein Rosenman and Gofman). Since in such instances the deposition of cholesterol probably is caused by local degenerative changes in the skin of the cyclids xanthelasma palipebrarum is best regarded as a localized lipoidous.

Histopathology The histologic changes are similar to those of primary xanthomatous. As a rule however fewer Touton cells are seen and they may be absent Fibrotic changes may occur (Mont gomery and Osterberg)

11 Aecrobiosis Lipoidica

This disease represents a localized lipoidosis masmuch as the deposition of lipids occurs in areas in which degeneration or necrobions of collagen has taken place. The necrobiosis of collagen is due to vascular changes (Roederer Woringer and Burgun). In those cases in which flubetes exists it can be assumed that the diabetes has caused the vascular changes. However diabetes originally thought to be a prerequisite for this disease is present in only about one third of the cases (Kaalund Jorgensen). In the other two thirds the Cause of the vascular changes is unknown.

Clinically one observes on the legs and rarely elsewhere on the skin one oxfeveral sharply demarrated irregularly outlined glared patthes which are yellow in the center and voloceous at the periph ery The center gradually becomes depressed and atrophic and may break down to form an uler.

Histopathology On histologic examination the eniderm . -

In these areas the collagen appears homogeneous swollen and partly basophilic (Fig. 132) The collagen bundles often are broken up and,

homogeneous material is present around some of the vessels and sweat glands

On straining with scarlet red a large amount of lipid material is visualized. It strains a rusty brown and occurs in the form of small droplets throughout the hyalin like material but particularly around the blood vessels (Fig. 131). It is located extracellularly On polari scopic examination the lipid material proves to be isotropic, not doubly refractile. Urbach and Wiethe interpreted the histologic findings as indicating a merging of lipid and protein and therefore suggested the name lipid proteinosis.

No definite conclusion has so far been reached about the future of the lipid substance in the dermis. In several cases, the chemical analysis of involved skin revealed a decided increase in the amount of lectiling which is a monorumophosphatide (Montgomery and Havens, Ramos e Silva). In other cases, however, no such increase was found (Wile and Snow, Hansen, Price LaRosa and Settle).

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Histopythology The involved skin shows a dense nonspecific cellular infiltrate composed mainly of <a href="https://linkinglight.org/linkinglight-public-linkinglight-p

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The cause of the disease is unknown Sobel and Pollock impressed

with the subnormal values for cholesterol in their is well is in Urbach's case suggest that for some unknown reason the blood is unable to hold normal amounts of cholesterol which therefore is

degeneration. In contrast with the neutral Fit in the subcutaneous layer and the sebaceous glands, which status a brilliant orange red, the graphlet statu a russi brown. According to Hildebrand, Mont

rol rol



Fig. 133 Secrobious lipoidica Several foreign body giant cells are located within an area of collagen degeneration. Two fibrotic vessels are present. (X200)

(Laymon and Fisher) The fact that occasionally, lipids are absent in the lesions (Sachs) indicates that their presence is purely a secondary phenomenon

translomators dischorn is chronica progressiva recently de scribed by several European authors (Viewher and Leder, Arzi) probably represents necrobiosis lipondica without lipid deposits

Differential Diagnosis. In the differential diagnosis, Lanuloma innulare must be considered, because both necrobiosis lipoulica and granulomi annulare show areas of collagen degeneration and the sine tipe of reactive inflammation and fibrosis. However, in granuloma annulare there are not associated that changes no deposis of lipids (ew or no grany cells and no atroph) or ulceration of the epidermis.

instead of lying parallel to the surface of the skin extend in various directions. There often is evidence of formation of young collagen between the degenerated bundles.

Within and near the areas of necrobiosis often extending into the subcutaneous fat one finds a predominantly perivascular inflamma tory infiltrate composed of symphocytes districtes dibroblasts and

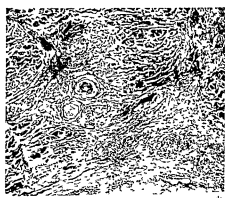


Fig. 132 Necrobiosis lipoidica. Much of the colligen appears de generated. An inflammatory infiltrate as scattered through the areas of degeneration. A vessel in the center shows endothelial prol feration and fibrosis of its vall. (×100).

occasional groups of epithelioid cells "Foreign body grant cells commonly are present and thus are of considerable diagnostic value (Michelson and Laymon Belote and Welton) (Fig. 133) Occasion ally a few form cells are noted (Klaber Nicholas)

The Glood vessels priticularly in the middle and the lower dermis exhibits fibrosis of their walls with proliferation of their endothelia lining. This process may lead to pritial and occasionally even to complete occlusion of the lumen. Thrombosis of small vessels occurs sometimes. These vascular changes account for the degeneration of

the collagen
Staining for fat with scarlet red frequently but not always reveals numerous granules of lipid extracellularly in the areas of collagen



Fig. 131 Primary systemic anyloidosis Round amorphous fissured masses of amyloid are present in the uppermost derms. They resemble close of colloid militari but in contrast to colloid militari are present throughout the dermis (X°00)

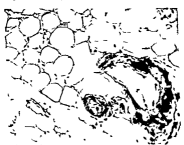


Fig. 13.5 Primary systemic amyloidosis Subcutaneous fat. Amy loid is deposited in the walls of two blood vessels and also around fat cells forming so-called amyloid rings (×400)

on the other hand, deposits of mucin usually are present. In spite of these differences, Ellis and Kirby Smith regard the two diseases as closely related.

AMYLOIDOSIS

Three forms of amyloidosis exist (1) primary systemic amyloidosis (2) primary localized amyloidosis of the skin (lichenoid amyloidosis) and (3) secondary systemic amyloidosis. In frimary systemic amyloidosis, the skin is frequently involved, invectondary systemic imploidosis, yery rarely

The Congo red test is of great value in the diagnosis of cutaneous amyloidosis. If 10 cc of a 15 per cent solution of Congo red is injected subcutaneously, or 01 cc intradermally, into the affected region, the areas in which amyloid is deposited will be standed strongly with red day after from 24 to 48 hours whereas the interposed skin will appear only very slightly stained (Nomland, Dostros sky and Sigher). In visiologic sections stained with we mutoxylin and conjuganyloid appears as 1

ruff etts, probably becau process of fixation. With the Yan Gieson stain, amyloid stains sellow and collagen red. Amyloid is a protein polysaccharide complex and therefore it stains at lenst in lichenoid amyloidosis and in secondary amyloidosis. Yed with Congo red. purple red with methyl volet (indicating metachromasia) and deeply red with the periodic acid schild reaction. In primary systemic amyloidosis, however, these staining reactions are not always present and warylfrom prinent to patient and even from organ to organ, probably because of differences in the amount of polysaccharides or in the nature of the union between the protein and the carbohydrate molecules (Goltz)

PRIMARY SYSTEMIC AMYLOIDOSIS

Mesenchamal Ussue is affected while the parenchyma of paren chymatous organs is spired. Amyloid deposits are found mainly in the smooth and striated musculriure in the <a href="mailto:smoothese:sm

Of interest is the frequency with which primary systemic amy loidosis is associated with multiple myelomi and Bence Jones proteimuria (Brunsting and MacDonald) Examination of the urine for Bence Jones protein, roentgenograms of the bones and sternal biopsy in search of atypical plasma cells in the bone marrow, therefore, should be performed in all cases of primary systemic amyloidosis

about capillaties (Nomland). In some cases, a mild chronic inflam

matory infiltrate will be found Differential Diagnosis Lichenoid amyloidosis must be differen

trated from colloid milium. For differential diagnosis see page 165

SECONDARY SYSTEMIC AMPLIOIDOSIS

This condition occurs in thronic suppurative diseases, such as tuberculosis and in thronic cachectic diseases associated with marked loss of protein from the body Amyloid deposits are found in the barenchymatous organs especially in the liner, the kidney, the spleen (sago spleen) and the adregats Thelpkin is unolled very rarely Michelson and Lynch have reported a case of secondary systems anylandosis due to tuberculosis in which there were diffuse nodular lesions of the lips due to deposits of amyloid

CALCINOSIS CUTIS

Two forms of calcinosis cutis exist metasiatic and metabolic calci fic ation

METASTATIC CALCIFICATION

Metastatic calcification develops as a result of hyperculcenia. The hypercalcemia may be due to parathyroid neoplasm hypersitaminosis D thronic renal disease or destruction of bone in such conditions as osteomyclitis and malignant growths (Mulligan) Colcium is and to be deposited in organs in which the cells excrete acid and therefore have a low carbon dioxide tension. As a result of the low and a

c cens or these organs excrete acid substances the skin and the subcutaneous tissue are affected only rarely. How ever instances of metastatic calcification of either skin or subcuta neons tissue have been reported as caused by paradistroid neoplasm (Penecle Laubmann) by hypervitaminosis D (Bevans and Taylor) by chronic renal disease (Platt and Owen) and by osteomyelitis (Weidman and Shaffer)

Histopathology Calcium deposits are recognized easily in histo four sections since they stain deeply blue with hematoxylin and cosin and blick with you kossy's stain for calcium. In most instances of inclustatic calculation, the calculm occurs as individual granules as well as massive deposits in the dermit and in the subcutaneous fat (Fig. 136) Larger deposits often evoke a foreign body reaction so that giant cells an inflammatory infiltrate and fibrosis may be present around them In Weidman and Shaffer's case the calcium de posits consisted of small granules which were found not only in the Clinically, the skin shows discrete and conlescing pipules and nodules, which are semitranslucent and of a way, amber color in addition, yellowish plaques resembling those of nanthomniosis are present. Petechiae and ecchymoses frequently occur at the site of eruption yitching is absent. The face is predominantly affected especially the periorbual regions.

Histopathology. Histologic examination of the skin reveals large faintly bosinophilic, amorphous masses of amyloid which may be deposited anywhere in the dermis (Fig. 134) as well as in the sub cutaneous tissue. In addition, small deposits of amyloid may occur in the membrana propria of the sweat glands around blood vessels and in the walls of blood vessels. In most cases, no inflammatory to action is present (Michelson and Lynch), but, in some cases, one find foci of lymphocytes, plasma cells and foreign body grant cells (Pear son, Rice and Dickens)

In the subcutaneous tissue, one may find besides large masses of amyloid and deposits of amyloid in the walls of the blood vessels so called amyloid arings which are formed by the deposition of amyloid around individual fat cells (Fig. 135) (Perison, Rice and Dickens). The fat cells thus appear as if cemented together by the cosmophilic amyloid substance (Iverson and Morrison).

Not only in the skin and the subcutaneous tissue, but also through out the body, the small arteries and the veins show amyloid deposits often entirely replacing their media and adventitia. The vessels of the tongue the skeletal muscle the respiratory tract, the heart and the gastro intestinal and the genito urmary tracts usually are affected most severely. In the skeletal muscle and in the tongue, numerous muscle fibers show amyloid deposits in the form of amorphous nodular swellings (Iverson and Morrison).

PRIMARY LOCALIZED AMYLOIDOSIS OF THE SKIN (LICHENOID AMYLOIDOSIS)

In this form of amyloidosis only the skin is involved. The lesions are seen most commonly on the legs but they may occur elsewhere. They consist of closely set discrete conical or flut, brownish red papules which resemble the pipules of hehen planus. Occasionally the pipules have a translucent appearance. In some instances the pipules, by coalescence form plaques which may develop a vertucous surface, and then resemble lichen simplex chronicus. The lesions usually tich severely in contrast with those of primary systemic amyloidosis.

Histopathology. The amyloid deposits are much smaller infine than those found in primary systemic amyloidosis and are limited to the subepidermal region of the dermis. The earliest deposits occur

In sclerodermic calcinosis, the histologic appearance of the skin and the subcutaneous tissue is that of scleroderma. The calcium deposits are usually located within areas of sclerotic collagen. The deposition of calcium probably is due to the decreased metribolic victivity in the sclerotic tissue. Decreased metabolic activity causes the carbon dioxide tension in the tissue to be lower than normal and thus reduces the solubility of calcium (Brodi and Bellin).

In patients with idiopathic calcinosis cutis no perceptible histologic changes may precede the deposition of the calcium (Fpitein Bruer Marble and Bennett Bauer, Marble and Bennett loand, in their case that the initial lesion consisted of deposition of finely divided particles of calcium salts around apparently normal far cells in the subcurancous issue. The granules seemed to coalexe slowly to form large masses. Other authors found thirt, even in idiopathic calcinosis, mild degenerative changes preceded the deposition of calcium (Rothstein and Welt Atlanson and Weber)

COUT

Court is a disturbance of purine metabolism characterized by <u>pathruis</u>. Deposits of urates are found in the tissues of the joints particularly their varializes in the cartilage of the ears and in the



Fig. 137 Gout Deposits of sodium bourate are surrounded by a foreign body grant cell reaction. On the left, the sodium hourate is present as needle shaped crystals. (X100)

276 Metabolic Diseases

dermis but also in the epidermis, in the sweat glands, in the sweat ducts and in the nerve trunks

METABOLIC CALCIFICATION

Metabolic calcification is due to local metabolic disturbances and is not associated with hypercalcenia. Deposits of calcium occur, as a rule, only in the skin and the subcutineous tissue, but occasionally



Fig. 136. Metastatic calcification as a result of hypercalcema (produced by prolonged and excessive idministration of virtumi D). Von kossa stam for cilcium. Irregular masses of calcium surrounded by a foreign body grant cell reaction are present in the subcut incous (i.t. $(\times 100)$).

also in the muscles and the tendons. The internal organs are spared. There may be just a few deposits (calcinosis circumscripia) or in numerable deposits everywhere in the skin (calcinosis universalis). Between 30 and 40 per cent of the cases of calcinosis circumscripia and universalis described in the literature occurred in pitients with seleroderima or dermatomyositis (Atkinson and Weber). (See 'Seleroderima,' page 310, and Dermatomyositis page 303.) In the rest, no reason for the calcinosis was evident.

Histopathology. As in metastric calcification, the calcium may be present as individual granules or as massive deposits in the dermis and in the subcutaneous fat. In the subcutaneous fat, extensive areas of calcification may be observed. A foreign body giant cell reaction is often found around the larger deposits of calcium.

In the delayed cutrunous form the bullae arise as pressure bullae (see page 66) subepidermally (Teligman and Baum Robert). However due to regeneration of the epiderms older bullae may be located partially or entirely within the epiderms. The bullae, thus, do not differ histologically from those observed in epidermolysis bullosa. The mila like those of epidermolysis bullosa dystrophica con see of small intradermally located epidermal cysis (Robert).

MYXEDEMA

Three types of myxedema occur generalized myxedema circum scribed myxedema and papular myxedema Generalized myxedema is a manifestation of hipothyroidism. Circumscribed myxedema is associated with or preceded by hyperhyroidism and occurs almost invariably together with exoplathalmos commonly it follows thy roidectomy or therapy with thiouracil. Papular myxedema is not associated with any disturbance of thyroid function.

The mucin present in the tissue in these three discuses appears light blue on staining with hematoxylin and eosin. It stains red with the periodic acid Schiff reaction and is strongly metachromatic with methylene blue thionine cresyl violet and toluidene blue indicating that it is a protein polysaccharide complex (Brewer). The mucin is digested in sections by hyldhronidase which points to the fact that it contains a large amount of hyduronic acid (Palitz and Brunner). It also stains red with mucicariume (The latter staining method requires fixition with absolute alcohol—see page 29).

GENERALIZED MUNEDEMA

Clinically the entire skin appears swollen dry pale and ways. It feels firm to the touch. In spite of its edematous appearance, the skin does not pit on pressure. The facies is characteristic, the nose is broad and thick and the lips are swollen.

Histopathology The dermis is increased in thickness. The collagen bundles as well as the individual fibers of the collagen bundles are separated by edema. The collagenous fibers show turns of decreases

o non mucin is usually small (Reuter)

CIRCL MISCRIBED MYXEDEMA

The lesions usually are limited to the anterior aspects of the legs. They consist of hard raised nodular yellow wavy plaques with prominent hair follicles.

subcutaneous tissue The deposits in the ears and in the skin mant fest themselves as nodules of varying size. These nodules are called tophi

Histopathology. The tophi show needle shaped crystals of sodium burrate lying closely packed in the form of bundles or shewes. The crystals often have a brownish color They stain well with you kossis stain. The accumulations of urates are surrounded by granulation tissue containing many foreign body giant cells (Fig. 137)

PORPHYRIA

Porphyria represents an inborn metabolic error in which large amounts of uroporphyrm and coproporphyrm are excreted in the urine. Three forms of porphyrm exist, the congenital form the acute intermittent form and the delayed cutaneous form. In the con genital form, the porphyrins are formed in the bone marrow (por phyria erythropoietica) while in the acute intermitten form and in the delayed cutaneous form they are formed in the liver (porphyria hepatica) (Watson)

In the congenital form, cutaneous lesions identical with those seen in hydrox vacciniforme appear from earliest childhood on the ex posed portions of the skin following exposure to the sin The lesions result in scarring and mutilation. The teeth may appear red

The acute intermittent form starts in adult life and is characterized by attacks of abdominal pain peripheral neuropathy and mental disturbances. Cutaneous lesions do not occur, as a rule. However, m rare instances, cut meous lesions like those in the delayed cutaneous form have been observed (Nesbut and Watkins)

The delayed cutaneous form (porphyria cutanea tarda) like the acute intermittent form, remains latent until adult life frequently chronic alcoholism by causing hepatic dysfunction precipitates the onset of clinical manifestations. Blisters form on exposure to light as well as on minor mechanical or thermal trauma. They often heal with scarring and formation of milia Abdominal pain and nervous system manifestations as in the acute intermittent type may be present, although to a lesser degree (Brunsting and Mason) Because of the occurrence of blisters following trauma and the presence of miles, some authors have referred to the lesions as acquired epider molysis bullosa. However, the exerction of uroporphyrm and copro porphyrm in the urine and the presence of sensitivity to light make it evident that the disease is not related to epidermolysis bullosi (Brunsting and Mason)

Histopathology In the congenital form the vesicles have the same appearance as in hydrox vacciniforme (see page 46)

In the delayed curmeous form the bullae arise as pressure bullae (see page 66) subepulermally (Jeligman and Baum Robert). However due to regeneration of the epidermis older bullae may be located partially or entirely within the epidermis. The bullae thus do not differ histologically from those observed in epidermolysis bullosa. The milia like those of epidermolysis bullosa dystrophica con sist of small intradermally located epidermal cysts (Robert).

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Histopathology. In the congenital form the vesicles have the same appearance as in hydrox vaccimiforme (see page 46)

PAPULAR MYXEDEMA (PAPULAR MUCINOSIS)

There is a widespread eruption of asymptomatic soft, yellowish pipules which may coalesce into irregular pritches.
Histopathology The appearance of the mucinous infiltrate is like that in circumscribed mysedema manmich as there are rather large. amounts of mucin as well as stell the cells (Dalton and Seidelf). However, the extent of the mucinous infiltrate is more limited than in circumscribed myxedema because it is present only in the upper dermis and in the case of individual papules within relatively small 31615

SCIEREDLMA ADULTORUM (BUSCHLE)

Scleredema adultorum is characterized by diffuse edema and induration of the skin and the subcutaneous tissue. Its cause is unknown but it is noteworthy that it frequently follows an infectious disease such as grippe or tonsillitis. It usually Legins on the face and spreads rishedly to intolve the neck and the upper trunk. Complete resolution takes place in a few month. The occurrence of pleural and peri cardial effusions and of hydrarthrosis has been reported (Vallee)

Histopathology Throughout the dermis one observes swelling and whiting up of the collagen bundles by edema. The edema may be severe enough to produce in the derinis clear unstained spaces of various sizes (fenestration) Freund noted that the edema sub state which did not stain with hemitoxylin and eosin stained meta chromatically with cress I violet like mucin but not with mucicar mude Vallee found this strining reaction in one case but not in mother Brum Falco observed purplish metachromasia with tolin dene blue which was no longer present when the sections prior to the staining were incubated with hyaluromidase. He concluded that the edema substance consisted of hyaluronic acid

AUDISON & DISEASE

Addison's disease which is caused by hypofunction of the adrenal clauds is characterized by weakness loss of weight low blood pres sure and diffuse hyperpagmentation of the skin and the mucous membranes

The hyperpigmentation in Addison's disease

of their is that in Addison's disease the dam aged adrenal gland responds but weakly to putuitary sumulation and

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Histopathology. Large amounts of mucin are present in the demnis particularly in the lower dermis. There it occurs not only as individual threads and granules, but also as massive deposits causing wide separation of the collagenous fibers (Fig. 138). The number of

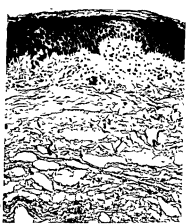


Fig. 138. Circumstribed myxedemi Considerable mounts of mucin are present especially in the lower derms separating the collagen bundles is well as individual collagen fishers. (The empty spaces are due to shrinkings and fulling out of mucin.) (X200)

fibroblasts is increased and newly formed collagen is present Some fibroblasts have a stellate shape and are surrounded by mucin. Be cause of the much larger amount of mucin and the new formation of collagen, the dermis is greatly thickened, much more than in generalized myxedema.

The epidermis shows hyperkeratosis. The rete ridges are often flattened. The vessels in the upper dermis are dilated and surrounded by a mild inflammatory infiltrate. Plastic tissue string show the elastic tissue to be frayed and greatly decreased (O Lerry).

Differential Diagnosis Differentiation of reanthosis nigricans from nerus verticosus may be impossible (Curth) As a rule however nerus verticosus shows more marked acuthosis thru acuthosis unput curton and occide elopment rather than atrophy of the rete ridges



Fig. 139 Acanthos's nigricans. There are hyperkerators and pap I lomators: Several pap liae project upvard as finger I ke project ons. As is usually the case acantl os's and hyperp gmentation are slight. (×100)

RIFHI S MELANOSIS MELANODERMATITIS TONICA POIKII ODERMA RETICULARE (CIVATTE)

Rights melanosis and melanodermatitis toxics have the same appearance clinically as well as histologically (Storck). In both conditions one observes ill defined symmetrical bluish brown hyperpigmentation of the face and occasionally also of the neck and the chest. There my be slight atrophy of the shin and follicular hyper-levitions. The cause of Rights dermatosis is not fully known but it has been suggested that it is caused by a lack of viramin B and provided by exposure to the sum and that this it is related to pellagra. Prehimann! Melanodermitis toxica is due to contact of the affected shin with vira oils or greases.

Polyhoderm reticulare differs from these two diseases clinically by its predominant localization on the neck, the reticular arrangement of the hyperpignetation and the presence of telanguertases (Phermi and Bosq). Histologically, the telanguectases usually are not sufficiently evident to differentiate polyhoderma reticulare from the other two diseases.

consequently, as a compensatory phenomenon, the pituitary gland is overactive.

Histopathology. Histologic examination often shows no changes other than hyperpigmentation. Occasionally, one observes slight flat tening of the rete ridges and slight thinning of the epidermis. The amount of melanin is increased in both the epidermis and the dermis. In the epidermis, the melanin is present chiefly in the bisal layer, but may be found also in the lower layers of the stratum indipglin Because the pigment formation proceeds slowly over a long period of time, the number of clear cells is increased only slightly. In the upper dermis, a moderate number of melanin laden chromitophores (melanophores) are present. There is no inflammatory reution in the dermis.

Differential Diagnosis. A diagnosis of Addison's disease cannot be made from histologic sections, because the same histologic picture is observed in nonspecific hyperpigmentation of the skin and in the normal skin of the Negro

ACANTHOSIS NIGRICANS

Three types of acanthosis nigricans exist mulignant acanthosis nigricans, benign acanthosis nigricans and pseudo teanthosis nigricans. Clinically and histologically, the three forms look ahke (Cnith)

The malignant form occurs in adults and is associated with in ternal cancer, usually of the glandular type. The benign or juvenile form may start at any time before pulberty. It represents a genodernatosis related to nevus vertucosus. Pseudo acanthosis nigricans occurs in the body creases of obese, brunette persons. It disappears when the patient loses weight.

Clinically, all three forms of acanthosis nigricans present verticous, hyperpigmented patches, predominantly in the avillae, on the neck

and in the submanimary and the genital regions

Histopathology. Histologic examination reveals marked hyperkeratosis and pupillomatosis. In addition, acanthosis and hyperpigmentation are present but are slight (Fig. 139). Thus, the name acanthosis

mericans has little histologic justification

In a typical lesion, the papillae project far upward as fuger like projections and are covered with a not unduly thickened straum malpighit. The valleys between the fuger like projections are filled in largely by keratin. In areas where there is no papillomatosis the epiderius shows areas of moderate acanthosis with adjacent areas of atrophy of the straum malpighit. The rete ridges, as a rule, are developed only poorly. There may be a slight increase in the amount of melanin in the basal layer.

VITILIGO

Viriligo is characterized by variously sized and shaped sharply demarcated patches of depigmentation surrounded by hyperpig mented skin

Histopathology In the depigmented areas the epidermis is devoid

ntain the

amount of melanin in the basal layer is increased and numerous

melanophores are present in the upper dermis

The dopa reaction is negative in the basal layer of the depigmented
areas and positive in the basal layer of the adjoining hyperpigmented
areas.

HEMOCHROMATOSIS (BRONZE DIABETES)

This disturbance of the metabolism is characterized by deposition of hemosiderin in various organs of the body. The presence of hemosiderin causes brouzing of the shin cirthosis of the liber and sclerosis of the spleen and the pancrers belerosis of the pancreas is responsible for the diabetes usually associated with the disease.

Clinically the pigmentation of the skin is diffuse and is indis

tinguishable from that seen in Addison's disease

Histopathology Granules of hemosiderin are found within chromatophores in the upper dermis and in the membrana propria of the wear glands Occasionally they may also occur in the basal cells of the epiderims. Hemosiderin is best demonstrated by staining sections with poissum ferrocyanide. With this method, hemosiderin granules on account of their content of iron, stain blue. Not infrequently melaini is present in excess in the basal layer and within chromatophores.

In selecting a site for biops, it is not necessary to choose a pig metid area because hemosiderin is present throughout the skin in hemosiderous. However it is important not to take a specimen from the legs where deposits of hemosiderin frequently occur in association with stass dermatitis and other vascular disturbances (Mont gomen and O Leary)

Differential Diagnosis It is impossible to differentiate the granules of hemosiderin from those of melanin in routine stains. They can however be easily differentiated by staining with potassium ferrocyanide which stains hemosiderin blue but does not stain melanin Furthermore melanin does not occur in the membrana propria of the sweat glands.

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Histopathology. The clinically visible hyperpigmentation in these three diseases is brought about by pigmentary incontinence of the bristl layer, resulting in an accumulation of melmin in the upper dermis. The epidermis shows mild hyperkeratosis, thinning of the stratum malpighii and varying degrees of degeneration of the cells in the basal layer. The amount of melmin in the birst layer is decreased. The papillary and the subpapillary layers of the dermis how

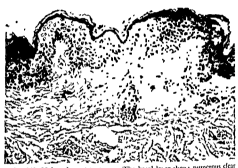


Fig. 110 Melanodermatuts toxica. The bisal layer shows numerous clear cells. The upper detrinis shows large amounts of melanin mainly within but also outside of chromatophores. A mild perivascular influmnatory in filtrate is present. (×200)

ever, show large amounts of melanin, mainly within, but also outside of, chromatophores (Fig. 140). In addition, there are various degrees of inflammatory reaction in the upper dermis. In some cases, the inflammatory infiltrate is limited to the perivascular areas, in other, it is extensive, brindlike and close to the epiderius, and thus resembles that of lichen planus (Storck).

At a later stage, the degeneration of the basal layer is slight and many clear cells may be present. The degree of inflammation lessens with the age of the lesion.

Differential Diagnosis Incontinentia pigmenti differs from the three diseases just discussed by showing no inflammatory infiltrate in the upper dermis. Addison's disease shows no inflammatory infiltrate either and only a slight amount of melanin in the dermis.

there may be considerable regeneration of the cutaneous appendages

(Stellens Bair and Sheard)

Differential Diagnosis Differentiation of phrynoderma from ich thiosis may be difficult. As a rule however richthyosis shows atrophy or absence of the granular layer thinning of the epidermis and elongation and branching of the rete ridges. Pityriasis rubra pilaris differs from phrynoderma by showing in addition to hyperkeratosis and follicular plugging spotted parakeratosis irregular acanthosis and a more pronounced inflammatory infiltrate not limited to the fair follicles.

PELLAGRA

Pellagra is crused by a deficiency of the vitamin B complex but particularly of mocutine acid. Besides cutaneous lesions pellagra usually presents also a stomatus which is characterized by edemined techness of the tongue and attophy of the lingual papillae. In addition gastro intestinal symptoms and nervous and mental changes may be present.

Cutaneous lesions occur predominantly on exposed areas such as the dorsa of the lands the wrists the face the Varea of the upper chest and the dorsa of the feet. In the early stage there is erythema which in severe cases may be accompanied by bullae. Later the erythema assumes a livid shade and the skin becomes thickned and scaling Ultimately the affected treas become atrophic and deeply permented.

Histopathology Early lessons present a chronic inflammatory infiltrate with moderate edema in the upper dermis. Vesicles and builde may be present. They may be located intra-epidermally as well as subepidermally.

Older lesions show hyperkeratosis with areas of parakeratosis and a moderate degree of acanthosis Follicular plugging occasionally is observed. The amount of melanin in the epidermis is increased. The

uceper portions of the dermis (Moore Spies and Cooper)

In the end stage hyperkeratosis and hyperpigmentation are still present but the stratum maipighii now shows considerable atrophy with flactering to the rete ridges. The dermis shows moderate fibrosis.

Differential Diagnosis The histologic picture of pellagra is not diagnostic. As a rule it is merely one of chronic dermatitis. In the end stage the presence of hyperpagmentation and of atrophy of the stratum malpighti serves to distinguish pellagra from chronic dermatitis.

The granules of silver, present in argyria, are also frequently located in the membrana propria of the sweat glands, but they differ from those of hemosiderin by being much smaller and more uniform in size Furthermore, they do not stain with potassium ferrogande and are refractife with dark field illumination (see page 153)

OCHRONOSIS

In ochronosis, due to an inhorn metabolic error, the carbolism of tyrosine cannot proceed beyond homogentisic acid. The disease is characterized by blackening of the cartilages and by osteo arthritis and may show bluish discoloration of the sclerae and brown of bluish mottled pigmentation of the skin. The turne darkens on exposure to the air. The discoloration of the cartilages, the sclerae and the skin as well as the darkening of the urine, are due to the presence of homogentisic acid, which by oxidation is converted into a dark-colored insoluble product.

Histopathology. Varying amounts of a light brown pigment are present in the dermis. The pigment is either diffusely distributed or present as clumps of varying size and shape. The clumps may be large, measuring more than 100 microns in diameter. The pigment does not stain with silver nitrate as melanin does but becomes black when stained with polychrome methylene blue (Laymon).

VITAMIN A DEFICIENCY (PHRYNODERMA)

Vitamin A plays an important part in the metabolism of the epithelial structures of the skin and the micons membranes. In addition to cutaneous changes, deficiency of vitamin A may cause night blind ness, xerophthalmia and keritomilacia.

The cutaneous changes, to which the name phrynoderma has been given, consist of dryness and roughness of the skin and the presence of follicular hyperkeratosis

Lichen spinulosus of Crocker is probably identical with phryno-

derma (Lehman and Rapaport)

Histopathology The skin shows modetate hyperkeratosis with marked distention of the upper part of the hur follicles by large horny plugs. In contrast with ichthyosis, the granular layer is present and may even be increased in thickness. The lower part of the livit follicles is atrophic and usually surrounded by a slight, chronic in flammatory inflirate. There are only few remnants of sebaceous glands. In addition, one my find evidence of atrophy of the sweat glands, such as flattening of the secretory cells (Frazier and Hu). In severe cases the sweat glands and the sebaceous glunds may undergo keratinizing metaplasia (Ressey and Wolbrich). When adequate amounts of vitamin A are supplied to a patient with phrynoderma,

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VITAMIN C DEFICIENCY (SCURY)

Scury, caused by a deficiency of ascorbic acid, is characterized by bleeding and spongs gums and petechial hemorrhages which often are perifollicular.

Histopathology. Scurvy is characterized by inability of the supporting tissues to produce and maintain intercellular substances (Wolbach and Howe) Thus, the endothelial cells of capillaries fail to form adequate amounts of intercellular ground substance Extravasation of red blood cells results. The extravasition occurs with out inflammators changes around the capillaries Scurys, conse quently, belongs to the group of noninflammatory purpuras (see page 127) (Peck, Rosenthal and Erf)

Histologic examination of the skin shows hemorrhages around the capillaries. As a rule, these hemorrhages are most pronounced in the vicinity of the hair follicles. In many instances, one finds hyperkera tosis and follicular plugging as in vitamin A deficiency According to Scheer and Keil, these changes in the epiderinis are purely secondary

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Systemic Diseases of Unknown Cause

1 UPUS ERYTHEMATOSUS

Three types of lupus erythematosus are generally terogramed (1) chronic discoud lupus erythematosus (2) subacute disseminate lupus erythematosus and (3) actue systeme lupus erythematosus (3) actue systeme lupus erythematosus and (4) actue lupus erythematosus and (5) actue lupus erythematosus (1) subacute lupus

type visceral lesions may be absent Inter

mediary forms between the three types occur chrically as well as littiologically. They cannot be regarded as separate diseases

Chaical Appearance Thronic discord lupus erythematosus is him tied usually to the face, where the flush areas of the cheeks are affected predomining. The scale and the eats also may be involved The testing course of well defined crysthematous slightly infiltrated patches showing adherent kerating cading and followlar plugging Older lesions show in addition atrophic scarring Systemic symptoms are shorter.

"Milacute disseminate lupus erythematosus hau may begin as such in udition to the fare other areas especially the thorax and the analysis and the arms and the legs, are moded Systemic symptoms such as lever malaise and legic peng, are other present. The levious consist of rythematosus others slightly livid patches which tend to coalesce and show may stight scaling. The prognosis as to life is fairly good since de scipment into the fuil a toute form of the discrete is not common to the substitute of the size of the size of the substitute of the su

Acture systemic lupus erythemytosus has severe systemic symptoms and almost invirably—Privil Fren corticotropin and cortisone as a rule metch delay death because the tern) lesions do not respond to these drugs. Systemic symptoms often precede the cutaneous eruption and occasionally curaneous manifestations are absent through out the course of the disease. The cutaneous eruption frequently begins on the face as a diffuse all-defined crythema with some edema.

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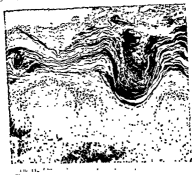
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fore, the hyperkeratosis may not be present until after the lesion is several weeks old Parakeratosis is usually completely absent. The keratotic filings are found mainly in the follicular openings but occur also in the sweat ducts and independent of either. The follicles inside the dermis may contain concentric layers of Leraum instead of hairs (Fig. 141). The atrophy of the stratum malpight is not always uni



There are . degeneration

form but may alternate with areas of acomhosis Focal liquefaction degeneration of the lasal layer represents the most significant histo long change in lupus crythematosus (Fig. 142). In its absence, a his rologic diagnosis of lupus crythematosus is rarely, if ever, justified in older lessons the basal layer often is partially absent as a consequence of his degeneration.

The sterms shows considerable edema in its upper portion. In drik's shanned persons utelanin is often present in the upper determs since, the degeneration of the cells in the basal layer causes them to lose their melanin (pigmentary incontinence) The capillaries and larger vessels are dilated and their walls may since deema, however,

("erythema perstans"). Soon other lesions appear, often widely spread over the body. They are poorly defined <u>purplish</u> and often edem tous, and may be <u>purpuric</u> or <u>vesicular</u>. The systemic symptoms include irregular leyer, malaise, weakness, pains in muscles and joints and pleural pain. Laboratory findings include marked leukopenia, hypergammaglobulinemia, proteinum and often, hematura



Fig. 141 Chronic discoid lupus erythematosus. Low magnification. There is scratour plugging and the follicles made the deemis contain, instead of hairs concentre layers of Jeratin. The epiderims is attoplise and devoid of referridges. The inflammatory infiltrate is distinctly patchy and tends to be located in the vicinity of hair follicles (X50).

-e -e ¹n most mbina m the

skin: (1) hyperkeratosis with keratotic plugging, (2) autopin of the stratum malpighii, (3) liquefaction degeneration of the basal cells, (4) a patchy, pervised by the concept of arrangement about degeneration of the collagen (11g 14t) Flowever, not all the changes

are present in every case.

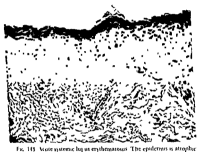
The pridermal changes are secondary to those in the dermis. There-

degeneration of the collagen (see under Acute Systemic Lupus Erythematosus) The of Acute Systemic Ludius Erythematosus The

. Warmord ration is observed

297

also in the skin. In order to study libranoid degeneration of the col



and shows marked fiquefaction degeneration of the basil layer. The dermis si ma fibrinoid degeneration. Only a mild permascular inflamma tory inflirate is present (×200)

lagen in the skin sections should be taken from covered areas of the body in exposed areas basophilic degeneration of the collegen --

ligen fibers i

mills not vis philic refract thickened ri was a sure those deeply cosmophilic than nor mills. In more advanced lesions the degenerated collagen fibers be come tragmented and have with the altered ground substance. The fibrinoid material shows metachromasia upon staining with toluidine blue and stains positive with the periodic acid Schiff reaction

proliferative or obliterative changes are absent. The inflammatory infiltrate is distinctly prachy. It is located mainly in the vicinity of the hair follicles and the seb-iccous glands, presses upon these structures and causes their gradual atrophy and disappearance (Fig. 141). The infiltrate is composed predominantly of lymphocytes but contains also a small number of plasmic cells and histocytes.

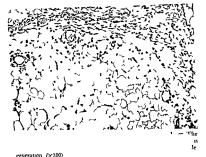
Unsophilic degeneration of the collagen in the upper derins is common in sections of discoid lupus erythemitosus obtined from exposed areas (where discoid fupus erythemitosus usually occurs) it is absent, however, in sections taken from covered areas (Mont gomery). In spite of its common occurrence in lupus erythematosus of exposed areas, bisophilic degeneration of the collagen cunnot be regarded in any way addiagnostic of lupus erythemitosus because it occurs also in simple senile atrophy of the exposed skin (see page 157) as well as in many other dermitoses when located in exposed areas. The clustic tissue shows, irrespective of whether the section is obtained from an exposed or a nonexposed area, at first fraging and later destruction throughout the dermis wherever the inflammatory infiltrate occurs.

In the differential diagnosis, two diseases have to be considered which share with chronic discoid lupus erythematosus the presence of a pitch; inditrate—namely, secondary sphilis and lymphocitic lymphoma Secondary syphilis differs from chronic discoid lupus erythematosus by the arrangement of the patchy infiltrate predominantly around blood ressels by the presence of numerous plasma cells in the infiltrate and by the presence of numerous plasma cells in the infiltrate and by the presence of vascular changes In lymphocytic lymphoma the patchy infiltrate is composed entirely of lymphocytes and does not show a tendency to arrangement near the epiderimal appendages. Neither secondary syphilis nor lymphocytic lymphoma show epidermal changes comparable with those of chronic discoid lupus crythematosus.

Histopathology of Subacute Disseminate Lupus Erythematosus
The Instologic changes in the lesions differ only in degree from those
of chronic discoid lupus erythematosus. The Introphy of the cpi
dermis, the Inquefaction degeneration of the bisal cells and the Idem
of the dermis are more prominent than in chronic discoid lupus
erythematosus whereas the hyperkeratosus and the Inflammatory in
filtrate are less marked

Occasionally, the edema in the upper dermis and the liquefaction of communication of the broad cells is severe enough to result in the formation of clefts and even vesseles between the epidermis and the dermis (McCreight and Montgomery) There may be evidence of fibringid

fibrillary ground substance normally not visible may be present as homogeneous eosinophilic fibrinoid clumps (Fig 143) (Klem perer Pollack and Baehr 1941) There is a rather mild perivascular infiltrate In addition one sees diffusely scattered instrocytes and fibroblasts some of which show pyknosis of their nuclei Extravasa tions of red blood cells are frequently observed Changes in the walls of the essels other than edema are usually absent. Only oc



generation (X100)

casionally does one see degenerative changes in the walls. They are hest demonstrated by the use of the periodic acid Schiff reaction (Stoughton and Wells)

The subcutaneous fat is often involved. It may show focal mucoid degeneration with reactive lymphocytic infiltration. The collagen bundles separating the fat lobules may be increased in thickness and show edema and fibrinoid degeneration similar to those in the dermis (Fig. 145)

Histopathology of the Visceral Lesions of Acute Systemic Lupus Erythematosus Visceral lesions usually are widespread but often minute in size so that they may easily be overlooked on gross inspection and even on histologic examination unless they are especially looked for The endocardium the serous membranes the heart and the skeletal muscle the renal glomeruli the spleen the lymph nodes and the fat depots are affected most commonly

Altshuler and Angevine believe that the fibrinoid material is formed by the precipitation of the acid mucopolysaccharides of the ground substance by an alkaline protein derived from either the necross of tissue or the interaction of the tissue with a damaging agent. The fibrinoid degeneration may affect the collagenous ground substance of capillaries arterioles and venules JThus viscular damage if pres

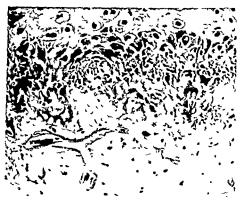


Fig. 144 Acute systemic lupus erythematosus. There are marked liquidation degeneration of the basil layer and utlema of the upper derm is. With not zone of subepidermal eddema, the collagen is present as homogeneous fibrinoid material which lies in irregular conglomerates and also surrounds a capillary (x400).

ent do s not represent in independent lesion but is part of the basic mury to the collegen (Kleinperer Pollack and Bacht 1941)

The histologic appearance of the skin in acute systemic lupus crythematosus resembles that of subacute disseminate lupus crythematosus (Madden Montgomery). One observes considerable liquefaction degeneration in the basal (ell layer and pronounced edema in the uppermost dermis (Fig. 143). Within the zone of subepidermal edema homogeneous cosinophilic fibrinoid material may be seen lying in irregular conglomerates as well as around crypilaries (Fig. 144). Deeper in the dermis some of the collagen bundles are swollen and stain intensely cosinophilic. The inter-

strated only very rarely in the blood of patients with diseases other than acute systemic lupus erythematosus a positive test for L. E. cells

is almost pathognomomic for it (Dubois)

The formation & E cells is due to the presence in the plasma of a factor which causes disintegration of the nuclei of neutrophils, as well as of other cells with subsequent phagocytosis of this material by neutrophils (Haserick, Gold) The L E cell is a neutrophil con



of a patient's blood after its incubation with heparin. In the left upper corner her an L. E rell a neutrophil containing a large smoky body In the right lower corner a rosette is seen consisting of amorphous ma terral surrounded by phagocytizing neutrophils (x400)

taining a round structureless smoky basophilic mass of such size that it presses the lobes of the nucleus against the cell membrane (Fig. 146) (Hargraves Richmond and Morton Smith) I. E. cells can be demonstrated most easily in smears made from the buffy coat of the patient's blood after its incubation with heparin (Dubois) Sinears of the buffy coat of heparinized sternal marrow may also be used. The smears are stained with Mright's stain. In addition to i. Firlis aread re n

- as and tosettes are precursors of the L. F. cell (Haserick)

The hematoxylin staining bodies are found in various organs, espe cially in the kidney and the endocardium but also occasionally in the

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The Yerrucous endocarditis of lupus erythematosus (the so-called Libman-Sacks syndrome) is caused by focal fibrancial degeneration of the subendothelial connective tissue of the endocardium and the subsequent round cell and fibroblastic proliferation and fibrosis Since fibrinoid degeneration occurs it in and again in the nealy formed fibrous tissue considerable autoints of granulomatous tissue are formed resulting in rused vertucous formations (I ibinin and Sacks Belgte and Ratner Gross Klemperer Pollack and Bachr 1941)

The scrous membranes such as the pleura the epicardium and the peritoned covering of the liver and the spleen may shot libri noid degeneration of their collegen with mild, reactive inflammation composed of hymphocytes, plasma cells distrocytes and fibroblasts

(Klempergt Pollack and Bachr 1941)

They avocardium and less frequently the skeletal muscle may show small foci of degeneration in the interfascicular connective tis sue and in the muscle bundles usually associated with a mild reactive inflammation. These changes are identical with those of demato. myositis though much milder (Klemperer Pollack and Bachr 1941 (ager)

The renal glomeruli not infrequently present wire loop lesions In this type of lesion individual glomerular loops appear thickened rigid and deeply cosmophilic because of fibrinoid degeneration and thickening of the basement membrane of glomerular capillates (Klemperer Pollack and Bachr 1911 Jager) In addition small foci of necrosis may occur in some of the glomeruli () ager)

Persarterial fibrosis of the central arteries in the wheen is one of the most common lesions in acute systemic lupus erythematosus Thick concentrically layered rings of sclerotic collagen fibers sur round these arteries (Klemperer Pollack and Buchr 1911 Kuser)
They mph nodes often show foci of necrosis

The body fit may show the same changes as those described for

the subcutaneous far

Vascular changes are usually not conspicuous Occasionally however the fibrinoid degeneration affects the collagenous ground substance of blood vessels. In thre instances, vascular changes are marked and resemble those of periateritis nodosi (Jarcho Malfors) it is possible that in these cases hipus erythematosus and periateritis nodosa are present simultaneously

Two structures are encountered frequently in acute systemic lupus erythematosus L E cells in the blood and hematowing saming bodies in the tissue They are absent in the food and hematowing saming disseminated lupus erythematosus Since L E cells can be demon strated only very rarely in the blood of patients with diseases other than acute systemic lupus erythematosus a positive test for 1. E. cells is almost pathognomogue for it (Dubois)

The formation M.L. E cells is due to the presence in the plasma of a factor which causes disuntegration of the nuclei of neutrophils, as well as of other cells with subsequent phagocytosis of this material be neutrophils (Haserick Gold) The L. F. cell is a neutrophil con



of a patient is blood after its metabation with heparin In the left upper corner hes an L. E. cell a neutrophil containing a large smoky body in the tight loner corner a roscue is seen constraine of amorphous ma terral surrounded by phagocytuning neutrophils (X400).

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The Yerrucous endocarditis of lupus crythemitosus (the so-called Labiman Sacks syndrome) is caused by focal fibrinoid degeneration of the subendothelial connective cususe of the endocardium and the subsequent round cell and fibroblastic proliferation and fibross. Since fibrinoid degeneration occurs again and again in the nealy formed fibrous cususe, considerable amounts of granulomatous tissue are formed resulting in rused verticous formations (Libiman and Sacks Beloie and Ratner Gross Klemperer Pollack and Bacht 1911)

The scrous membranes such as the pleura the epicardium and the peritoneal covering of the liver and the spleen may show fibrated degeneration of their collagen with mild, reactive inflammation composed of lymphocytes, plasma cells histocytes and fibroblass (Klemperet Polital, and Baeir 1941)

(Klemperet Politick and Brehr 1941)
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Periarterial fibrosis of the central arteries in the spleen is one of the most common lesions in acute systemic lupus erythemitions. Thick concentrically layered rings of sclerotic collagen fibers surround these arteries (klemperer Pollack and Baehr 1911 kniser).

The simple nodes often show foce of necrosis

The body fat may show the same changes as those described for the subcutaneous fat

Vascular changes are usually not conspicuous Occasionally however the fibrinoid degeneration affects the collagenous ground substance of blood vessels. In the manners assemble thinges are marked and resemble those of permateritis modosa (Jaicho Mallory). It is possible that in these cases lupus crythematosus and permaterius modosa are present simultaneously.

"The structures are encountered frequently in acute systemic lipus erythematosus L E cells in the blood and hematoxylingtuning bodies in the tissue. They are absent inchronic discoil and chronic discoil and

LUPUS ERYTHEMATOSUS PROFUNDUS (KAPOSI IRGANC)

In this rare condition (utaneous lesions of chronic discoid lupus crythematosus are present and in addition one or more firm sharply outlined movable <u>subcutaneous nodes covered</u> by normal appearing skın

> s that because subcurreous

sarcond of Darier Roussy He concludes that lupus erythematosus profundus constitutes a coexistence of two different diseases-discoid lupus erythematosus and subcutaneous sarcoid of Darier Rouss

DERMATOMYOSITIS

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In dermatomyositis the skin and the skeletal muscles are pre dominantly affected

The cutaneous lesions consist of extensive rather sharply defined areas of erythema and edema involving predominantly the face the chest and the arms. The eruption often greatly resembles that of subacute or acute lupus erythematosus. The lesions in the muscles cause progressive weakness with vigue muscular pain and later atrophy of the muscles Involvement of the esophagus often results in dysphagia Prior to the use of corticotropin or cortisone the dis ease was often fatal. Even with its use atrophy and fibrosis of the skin and the muscles may result producing a clinical picture re sembling that of generalized scleroderma

Histopathology In the skin particularly in early lesions the his tologic changes may resemble those of subacute disseminate lumis erithematosus (kinney and Maher O Leary and Waisman) Not in frequently however the histologic picture is that of a nonspecific chronic dermatitis. In old lesions, the collagen bundles of the dermis may show thickening homogenuation and selerosis and the cuta neous vessels fibrotic thickening of their walls so that the changes are indistinguishable from those of scleroderma (Dowling Freuden (led)

The subcutaneous fat may show mucoid degeneration of the fat cells and focal lymphocytic infiltration in the early stage and areas of fibrosis and calcification in the late stage (Norregaard Wainger and Lever) (see Calcinosis Cutis page 278)

During the active phase of the disease the skeletal muscles show

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skin (Klemperer, Gueft, Lee, Leuchtenberger and Pollister, Gueft)
They appear in sections stained with hematoxylin and eosin as red
purple, homogeneous bodies of the size and the shipe of fibroblast
or histiocytes They may be present singly or in aggregates They de
velop from degenerating nuclei and are identical with the smoly
body within L. I. cells Occasionally, one may see histiocytes or
neutrophils which have phagocytized a hematoxylin staining body.
The similarity of these cells to L. F. cells is striking. Histochemical
investigations have shown that, in the smoly bodies of L. E. cells
and in the hematoxylin staining bodies, the Feulgen reaction for
desoxyribonucleic acid (DNA) is positive. This is evidence that they

are derived from nuclear chromatin (Gueft)
Relationship of Acute Systemic Lupus Erythematosus to Der matomyositis and Generalized Scleroderma. These three disease—
lupus erythematosus, dermatomyositis and scleroderma—are related to one another (Banks). The fundamental pathologic lesion, namely fibrinoid degeneration of the collagen, is the same in all three diseases (Klemperer, Pollack and Baehr, 1941, Baehr and Pollack). Only the response to the fundamental lesion and the organs affected differ in these three diseases. One may summarize the response to fibrinoid degeneration of the collagen in these three diseases as follows.

In lupus erythematosus, the degeneration is associated with mild inflammation, occurring, as a rule, in small foci. The lesions are widespread

In dermatomyositis, the degeneration is associated with usually pronounced inflammation. Lesions are present mainly in the skin and the striated musculature, but other organs, particularly the heart may be involved.

In scleroderma the degeneration is associated with only slight in flammation but with marked fibrosis. As in dermitomyositis the principal lesions are in the skin and the striated musculature. In many cases, other organs (such as the heart, the esophagus, the lungs and the kidneys) are involved also.

The cause of fibrinoid degeneration of the collagen in these three diseases is not known. It is assumed by some that the degeneration represents an allergic or hyperergic, reaction to infection, particularly streptococcal infection. However, Klemperer, Pollack and Baehr (1942) point out that, although fibrinoid degeneration of the collagen may occur in hypersensitivity rections, it does not necessarily follow that all instances of fibrinoid degeneration are on the basis of hypersensitivity.

In older lesions the changes in the muscles may resemble those of scleroderma. The muscle bundles show sclerosis and utrophy and scieroderma. The muscle bundles show scierous and trophy and fibrotic connective tissue replaces the muscle bundles in many areas. On the basis of a study of these lite changes as just described in the skin and the skeletal muscle several authors have concluded that dermatomyositis and generalized scleroderma represent one and the same disease (Dowling Freudenthal)

Changes in organs other than the skin and the skeleral muscles

occur but not so regularly as in acute systemic lupus crythemitorus or in generalized scleroderma. The heart may show changes identi cal with those in the skeletal muscle though less severe (kinney and Maher O Leary and Waisman Wunger and Lever) The body fat may be affected similarly as the subcutaneous fat (Greenway and Lambie Kunney and Maher Wainger and Lever) Inflaminatory changes in the serous membranes may occur (kinney and Maher)
I kerative Jesions in the gastro intestinal tract due to vascular occlu sions have been described (Karelitz and Welt Horn Wringer and Lever)

Of interest is the relatively common occurrence of visceral carci noma in patients with dermatomyositis (Dostroisky and Sigher) So far 31 cases of carcinoma in dermatomyositis have been reported (Schuermann) In a statistical analysis Schuermann concludes that dermatomy in a statistical appayers Schiedmann continues that dermatomyosius than in the normal population. In contrast with this only one case of carcinoma in scleroderma has ever been reported

POIKILODERVIA ATROPHICANS VASCULARE (JACOBI)

Poikiloderma atrophicans vasculare has been described in a few instances as an independent disease (Marchionini and Rosser Dow

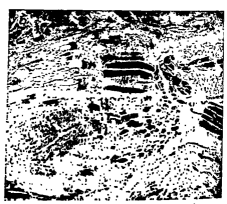
> As a rule It prouts

IN OCCUIV and in micross fungoides (see page 481). When associated with der matemyosius the term poishlodermatemyosius is often employed. Clinically poishloderma atrophicans assculare presents large ill defined areas usually in symmetrical distribution, which in the early

stage show erythema and slight scaling a mouled

o e promounced The clinical picture then resembles chronic radiodermatitis

degenerative changes and inflammation. The degree of these changes varies not only in different muscles but also within each affected mus cle. Even in severely affected muscles, close to areas of severe damage, one often finds areas of mild degeneration and areas in which the muscle bundles appear normal (Fig. 147). In areas of mild degenera tion, the muscle bundles exhibit effacement of transverse striction,



Γιο 117 Dermatomyositis, muscle The muscle bundles show viri ous degrees of degeneration. In addition, one sees edema and focal collections of inflammitory cells (×100)

coagulation or hyalinization of the sarcoplasm and proliferation of nuclei. In more severely degenerated areas, the muscle bundles show fragmentation of fibers, granular and vacuolar degeneration, baso philic staining and phagocytosis by large histiocytes. Inflammatory changes are secondary and not essential; they represent a reaction to the parenchymatous damage. One observes a cellular infiltrate, com posed largely of lymphocytes, but containing also plasma cells, his tiocytes and fibroblasts, between the muscle bundles, either in peri vascular arrangement or distributed diffusely. In addition, edema usually separates the muscle bundles The blood vessels are dilated, but, as a rule, their walls reveal no abnormalities (O'Lear) and Waisman).

In older lesions the changes in the muscles may resemble those of scleroderma. The muscle bundles show sclerosis and atrophy and scierogerma. The muscle bundles show scierous and incoping and fibrotic connective tissue replaces the muscle bundles in many areas. On the basis of a study of these late chinges as just described in the skin and the skeletal muscle several authors have concluded that dermatomyositis and generalized scleroderma represent one and the

same disease (Dowling Freudenthal)

Changes in organs other than the skin and the skeletal muscles occur but not so regularly as in acute systemic lupus etythematosus or in generalized scleroderma. The heart may show changes identi or in generalized scleroderim. The heart may snow changes identical with those in the skeletal muscle though less severe (kinney and Waher O Leary and Wasiman. Wringer and Lever). The body furmay be affected similarly as the subcutaneous fat (Greenway and Lambie kinney and Maher. Wainger and Lever). Inflammatory changes in the serous membranes may occur (kinney, and Maher). Ulcerative lesions in the gastro-intestinal tract due to vascular occlu. sions have been described (Karelitz and Welt Horn Wainger and Levery

Of interest is the relatively common occurrence of visceral earch noma in patients with dermatomyositis (Dostroisk), and Sighter). So far 31 cases of carcinoma in dermatomyositis have been reported (Schuermann). In a statistical analysis. Schuermann concludes that (Stituchmann) In a manifest analysis occurring that continues that carcinoma occurs at least five times more frequently in patients with dermatomyosius than in the normal population. In contrast with this only one case of carcinoma in scleroderma has ever been reported

POINTLODERMA ATROPHICANS VASCULARE (JACOBI) Poikiloderma atrophicans vasculare has been deserted

most frequently, in dermatomyouts but also in lupus crythematosus and in mycoss fungoides (see page 484). When associated with der matomyouts the term poskholdermatomyouts is often employed. Clinically poskholderma atrophicans vasculate presents large all defined areas usually in symmetrical distribution which in the early stage show crythema and slight scaling a mottled pigmentation and numerous telangectaies. In the late stage the skin appears atrophic the crythema has lyigely disappeared but the mottled pigmentation and the telangectains are more pronounced. The clinical picture twen resembles chronic radiodermatics. then resembles chronic radiodermatus

Histopathology. In the idiopathic form of poikiloderma and in poikiloderma associated with derinationyosits or lupus explient tosus the histologic changes are identical. In the early active stage the epidermis shows moderate atrophy of the stratum malpighic effecience of the rete ridges and hydropic degeneration of the basicells (Fig. 148). In the upper dermis one finds a furly dense cellular



Fig. 148 Poikilodermatomyositis early stage. The epiderms sloss atrophy and hydropic degeneration of the basal cells. In the upper dermis one sees a furily dense inflammatory infiltrate, which in places invades the epidermis. The collagen of the upper dermis shots edema and hyalimization (x200).

infiltrate which in places invides the epidermis and often has a band like arrangement. The infiltrate consists of lymphocytes histocytes and fibroblasts. Many inclanophores may be present. Some of the superficial capillrates are diluted. The collagen is edematous and shows hydine degeneration. The clustic tissue is largely destroyed. Hur follicles and sebucious glands, are absent. (Horn)

In the late stage the epidermis is attoplic. There is marked dilation of superficial capillaties. The dermis shows homogenization and sclerosis of the collagen with little or no inflammatory inflictive Melanophores are present in varying numbers. In contrast with scleroderma, the dermis is greatly thinned (Guy Grauer and Jacob)

In polkhoderma associated with mycosis fungoides the histologic changes in the dermis are those of mycosis fungoides (see page 489)

Differential Diagnosis Subacute disseminate lupus crythematosus like poikiloderma attophicans visculare shows attophy of the stra tum malpoginu and vacuolitation of the basal layer. However the presence of the superficially located brindlike infiltrate in poikilo dermi attophicans vasculare usually makes a differentiation possible.

SCLERODERMA

Two forms of scleroderms occur circumscribed scleroderma (mor plical) and systemic or generalized scleroderma

Clinical Appearance In circumscribed scleroderma (morphea) one or several round oral or irregularly shaped smooth indurined parches are present. They are at first dull red or violucous in color but soon resume an ivory color. As long as there is peripheral extension the patches tend to have a purplish Irilo (lilac ring.) The disease is beingly fluterial lessions are absent.

In systemic or generalized scieroderma, large areas of the skin are affected. At first, the involved areas present diffuse induration. As the disorder progresses, the skin and the subcutameous tissue become firmly bound to the underlying structures so that motion is difficult. Gradually, the skin and the subcutaneous tissue undergo strophy but even at this stage the skin retains some of its induration. The face and the hands are often the most severely affected areas (acroscleosis). In addition to the skin the strated musculature naturably is affected resulting in weakness and muscular atrophy. Involvement of the esophagus may lead to difficulties in deglisition of the first to cardiax insufficiency of the lungs to dyspore.

Histopathology of Circumscribed Scieroderma (Morphea) An early inflammatory and a late scierone stage exist. Most sections obtained routinely show a histologic picture intermediary between the stages.

In the early lesion the collagenous bundles appear swollen and homogeneous. They are separated by edema. An inflammatory inflittate predominantly lymphocytic is present between the collagenous bundles and around blood vessels (Fig. 149). The walls of the vessels are edematous. These changes are present throughout the dermis In addition the inflammatory inflitrate extends between the fat cells of the subcutaneous layer causing degeneration of the fat cells. The classic fibers are trajed, and may be destroyed.

In a fate lesson the dermin is markedly thickened. The collagen bundles are hypertrophic selectotic and closely packed (Fig. 150) Hirrobitus are fewer than in the normal dermis. The inflammatory infilitate has disappeared almost completely, except around the vessels where as a rule one still finds a few inflammatory cells. Most of

the vessels in the dermis show marked thickening and sclerosis of their walls with narrowing of their lumin? Sebaceous glands and hair structures are often completely absent Sweat glands on the other hand, are still present, they are reduced in number and are atrophic Instead of lying close to the cutaneous subcutaneous border and being surrounded by fat cells they he in the midst of sclerotic

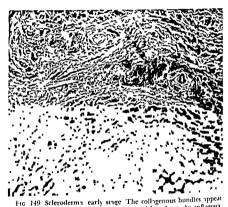


Fig. 149 Scierodermi early stage. The Configuration should and homogeneous and are separated by edema. In influence tory inflitrate predominantly lymphocytic is present between the collagenous bundles. The wall of the blood vessel seen on the right side is thickened by edema and beginning fibrosis. (X100)

collagenous bundles tightly bound down by them The fact that the sweat glands he inside the dermis rather than at its lover border is evidence that the thickening of the dermis is produced not alone formation of collagen at the lower border of the dermis. The border between the dermis and the subcutaneous layer is not so sharp as it usually is because thic strands of sclerotic collagen extend from the dermis into the subcutaneous layer may be greatly attended. The thickness of the subcutaneous layer may be greatly reduced. The vessels in the subcutaneous layer, including those of large caliber, often show marked thickning of all their coats with

natrowing of the lumen (O Leary and Nomland) It is worth noting that the rete ridges of the epidermis usually remain well preserved in spite of the fact that the thickened collagen bundles extend right up to it

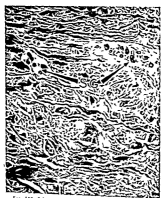


Fig 150 Scleroderma late stage The collagen bundles

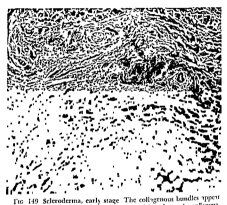
(×100) (×100) to / Mr tigitly bound down by collagen

Histopathology of Generalized Scienderma. The cutaneous that a histologic differentiation of the two types is not room between the cardy stage.

the collagen

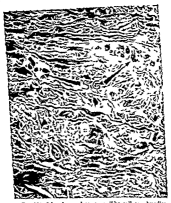
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the vessels in the dermis show marked thickening and sclerosis of their walls with narrowing of their lumina. Sebaceous glands and hair structures are often completely absent. Sweat glands, on the other hand, are still present, they are reduced in number and are atroplic. Instead of lying close to the cutaneous subcutaneous border and being surrounded by fat cells, they he in the midst of sclerote.



Tie 149 Scleroterma, early stage The Configuration between a smollen and homogeneous and are separated by edemy An influent tory infiltrate, predomin intly lymphocytic, is present between the col lagenous bundles. The will of the blood vessel seen on the right side is thickened by edema and beginning fibrosis (×100)

collagenous bundles, tightly "bound down' by them The fact that the sweat glands he inside the dermis rather than at its lower border is evidence that the thickening of the dermis is produced not alone by hypertrophy of the pre-visting collagen bundles but also by new formation of collagen at the lower border of the dermis. The border between the dermis and the subcutaneous layer is not so sharp as it usually is, because thick strands of sclerotic collagen extend from the dermis into the subcutaneous layer and replace much of the subcutaneous fat. The thickness of the subcutaneous layer may be greatly reduced. The vessels in the subcutaneous layer, including those of large caliber, often show marked thickning of all their coats with



F.c. 150 Scleroderma late stage: The collagen bundles are hypertroph c sclerot cand closely packed together. Othy vary I tile inflamm story sublinate as precent Fairoblasts are fever than in the normal desims: Groups of rather attophics eat glands (S.G.) are 1 ghthy bound do n by collagen (x100).

Histopathology of Generalized Scleroderma. The cutaneous changes are eigenfully the same as in circumscribed scleroderma so that a histologic differentiation of the tino types is not possible. In the early stage degenerative changes are often more severe than in the circumscribed type. Doe may observe fibrining degeneration of the collagen (Pollack). The walls of the vessels may show marked intend prohiferation fit mould degeneration and inflammatory in filtration (Massug and Va Shu). An occasional vessel may show throm

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bosis. In the late stage, focal or even extensive calcification may take place in the lower dermis and in the subcutaneous layer (Talbott Gall, Consolazio and Coombs, Brody and Bellin, Kanee) (See also "Calcinosis Cutis," page 276) Also, in the late stage of generalized scleroderma, the epidermis may show, in contrast to circumscribed scleroderma, atrophy with disappearance of the rete ridges

Histopathology of the Visceral Lesions of Generalized Scleroderma. There are often extensive systemic manifestations The skele tal muscles are nearly always affected severely. In early lesions de generation of muscle bundles with accompanying inflammation similar to dermatomyositis but less severe, may be observed occa sionally However, the characteristic changes are homogenization sclerosis and atrophy of the muscle bundles with increase in the in terstitial connective tissue. In contrast with dermatomyositis, the interseptal blood vessels often show marked obliterative changes Foci of chronic inflammation may be present around the vessels. The musculature of the esophagus is often affected similarly (Lindsa), Templeton and Rothman) Extensive atrophy and fibrosis may occur in the heart muscle (Weiss, Stead, Warren and Bailey) and in the intestinal musculature (Bevans) Widespread sclerosis of the medium sized and the small sized vessels in the myocardium has been ob served

Pathologic changes may occur also in the endocardium and the epicardium (Pollack), in the serious membranes (Bevans) and in the esophageal and the intestinal mucosa (Bevans). The changes consist in early lesions, of fibrinoid degeneration of the collagen with reactive inflamination, as in acute systemic lupus erythematosus. In older lesions, homogenization and sclerosis of the collagen pre-dominate.

Additional findings in occasional instances include glomerulitis of the wire loop type, as in acute systemic lupus erythematosus (Pollack, Bevans) extensive fibrosis with cystic changes in the lungs (Dostroysky), and fibrosis of the thyroid (Bevans)

PERIARTERITIS NODOSA

Periarteritis nodosa is a minifestition of hypersensitivity. All though, in most cases, the cause of the disease is not apparent, the administration of foreign serum or of sulfonamides may be the cause (Rich). The disease affects mainly small arteries and arterioles, but occasionally also veins. The principal vessels affected are those of the gastro intestinal tract, the kidneys and the heart, but those of the brain, the lungs and the skin as well as those of other organs may be involved.

Depending on the sites of involvement the clinical symptoms may vary Severe abdominal pain and symptoms of nephristis and of myocardial disease are the most common manifestations. Irregular lever and marked prostration are present. The disease is fatal in the vast majority of cases.

Cutaneous manifestations are found in about one third of the cases. They are manifold and may consist of macules papilles nod utes perechiae ecchymoses and necrotic uters (Fig. 151). Subcutaneous nodules usually movable and painless are observed occasionally (ketron and Bernstein). In some cases, extensive areas of cutaneous hemorrhage with subsequent necrosis occur (Melczer and Venker).

Histopathology The name periarteritis is misleading for the lesions actually represent a panarieritis. On a histologic basis the changes which occur in the arteries and the arterioles may be divided into four stages (Arkin Weir).

In the first, the degenerative stage, sections of the intum; and the media undergo nectors. The necrosis often affects only a segment rather than the entire circumference of the vessel, and only portions of the vessels are involved. Military meanysms may form in areas of segmental necrosis.

In the second the inflammatory stage the necroire area and the adjoining adventura are densely infiltrated with polymorphomelear leukocytes and cosmophils and sometimes also with lymphocytes and plasma cells (Fig. 152). The infiltrate extends to the permaneular tissue. The lumen of most vessels thus affected is thrombosed.

In the third the granulation stage, the necrotic part of the vascular wall is replaced by granulation tissue and the intima shows probleration leading to partial or total occlusion of the lumen

In the fourth the fibrotic stage the destroyed vascular wall is re placed by serr tissue. The lumen may show reduction in size obliter attom or recanalization.

In the skin the arterioles at the cutaneous subcutaneous border and in the subcutaneous tissue may show the typical changes of periorterius nodosa (Fig. 151). However, the small blood is

phils and eosinophils man) of which show fragmentation of their nuclei Thrombus formation with or without inflammatory reaction also may occur in the cutaneous ressels. The vasculitis and the throm bus formation cause extravasation of erythrocytes (Ketron and Bern stein).

It has been noted by several authors (Carol and Prakken, Miescher, Slinger and Starck) that periarteritis nodosa may occur in a bengn,



Fig. 151 Periarteritis nodosa, Low magnification An artery (A) at the cutaneous subcutrineous border shows the granulation stage of periarteritis nodosa. Above the artery is an ulcer which is probably caused by the occlusion of the artery (x50)

chronic form limited to the skin and the subcutaneous tissue. These authors state that, histologically, the lesions of the benign form are indistinguishable from those of the malignant systemic form.

ALLERGIC GRANULOMATOSIS

An attempt has been made recently to divide periarteritis nodosa into two types: "true periarteritis nodosa" and "allergic granulo matosis" (Churg and Strauss, Strauss, Churg and Zak; Zeek) In the

latter group allergic manifestations dominate the clinical picture to a much greater extent than in true pertacteritis nodosa. Most pratients have severe asthma as the first symptom and in addition to tients have severe asthma as the first symptom and in addition to the product of the contact of the conta

pronounced The course is

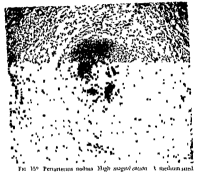


Fig. 15° Persarterius nodosa High magnit cateda. V thedium-sized artery located in the lower derinis shows partial nectors of its vall and invasion by inflammatory cells (x200).

chronic and usually fatal. Strauss. Ching and Zak believe that periarteritis nodosa as a result of dring sensitivity usually manifests itself as allergic granulomatous.

Histopathology Histologically in addition to especial widespread lessons of pernatterius nodosa one finals extratascular granusomas in many organs. The cutaneous and the subcutaneous nodules show no necrotizing atteritis but only extratascular granusomas. They consist of areas of central necrosis surrounded by radially airranged instrucytes and foreign body grant cells which are embedded in a diffue inflammatory inhitrate rich in cosinophilis (Strauss Churg and Zak).

The purpure lesions show the same appearance as those of ana phylactoid purpura (see page 127)

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bus formation cause extravasation of erythrocytes (Ketron and Bern stein).

It has been noted by several authors (Carol and Prakken, Miescher, Slinger and Starck) that periarteritis nodosa may occur in a benign



Fig. 151 Periarterits nodosa Low magnification. An artery (4) at the cutaneous subcutaneous border shows the granulation stage of periarteritis nodosa. Above the artery is an ulcer which is probably caused by the occlusion of the arters (x50).

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Epidermal Tumors

HISTOGENESIS AND CLASSIFICATION OF EPIDERMAL THMORS

The epidermal tumors may be divided into two classes namely tumors of the surface epidermis and tumors of the epidermal append ages In each class nevoid tumots (or hamartomas) and caremomas OCCUL

Second tumors or hamariomas are defined as beingh neoplasms which usually arise in accordance with the Cohnheim theory from may arise from immature

ring adult life Carcinomas

posed according to Hanse mann's theory of anaplastic cells. They usually arise from mature cells due to their change into anaplastic cells but embryonal cells

+ it all autonomous

able to exist more independently than embryonal cells and there fore survive when carried away through the lymphatics and multiply as metastases

Tumors of the Surface Epidermis The tumors of the surface epidermis may be classified as follows

- 1 Nevoid (benign) tumors
 - a Nevus verrucosus (papilloma) b Fpidermal con
 - 9 Precancerous tumors
 - a keratoris semilis
 - b Leukoplakia 5 Carcinomas
 - a Squamous-cell carcinoma (epidermoid carcinoma)
 - b Bowen's d sease (nira ep dermal squamous-cell care norma) c. Paget s disease

Tumors of the Epidermal Appendages The nevoid tumors or hamattomas of this group can be divided according to the decreas ing degree of organization and differentiation observed in them into

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Epidermal Tumors

HISTOGENESIS AND CLASSIFICATION OF EPIDERMAL TUMORS

The epidermal tumors may be divided into two classes, namely, tumors of the surface epidermis and tumors of the epidermal append ages In each class nevoid tumors (or hamariomas) and carcinomas occur

Nevoid tumors or hamartomas are defined as benign neoplasms which usually arise in accordance with the Colinheim theory, from

mann's theory of anaplastic cells. They usually arise from mature cells due to their change into anaplastic cells, but embryonal cells may also occasionally change into anaplastic cells and thus give rise to carcinoma. Anaplastic cells are dedifferentiated cells which behave differently from embryonal cells (Foulds). They are autonomous, able to exist more independently than embryonal cells and, there fore survive when carried away through the lymphatics and multiply as metastases

Tumors of the Surface Epidermis The tumors of the surface epidermis may be classified as follows

- 1 Nevaid (benign) tumors
 - a Nevus verrutosus (papilloma)
 - b Epidermal cist 2 Precancerous tumors
 - a Keratosis senilu
 - b Leukoplakia 3 Caremomas

 - a Squamous-cell carcinoma (epidermoid carcinoma)
 - b Bowen's disease (intraepidermal squamous cell carcinoma) c. Paget a disease

Tumors of the Epidermal Appendages The nevoid tumors, or hamartomas of this group can be divided, according to the decreas ing degree of organization and differentiation observed in them, into

Epidermal Tumors

four groups organic next adenomis beingin epithelioms and basi-cell epitheliomis (See Table 6). The carcinomas of the epidermal appendages can be divided into sebaceous gland carcinomis eccine gland carcinomis and apocine gland carcinomis. The author has advanced the Miesis that the organic next the adenomias the beingin epitheliomis and the brisal cell epitheliomis develop from arrested embryonal primary epithelial germ cells and, as such are primary epithelial germ tumors. Since the primary epithelial germ is an embryonal structure (see Chart I page 4) the tumors developing from it can accordingly be regarded as nevoid tumors or humarrooms. It is the author's belief that the organic next the adenomic and the beautiful combining arise from primary cells. the adenomas and the benign epitheliomis arise from primary epi theirs germ cells which prior to the onset of neoplasia ritained a certrun degree of differentiation whereas the basil cell epitheliomas arise from primary epithelial germ cells which attrined no or only title differentiation in accordance with the potentialty inherent in the primary epithelmi germ to differentiate into sebaceous glanda apocrine glands and hair differentiation in the timors developing from the primary epithelmi germ can be toward either sebaceous glanda apocrine gland or hair structures (see Table 6). Pinkus regland sporting gland or hair structures (see Table 6). Pinkus recently has suggested that those tumors in this group which arise in luter life—especially the basal cell epitheliomas—do not necessarily arise from congenitally preformed epithelial germ resis but from imprature pluripotential cells forming in later life. Whe origing her i are composed of matture organic structures. The currencous adenomias show less differentiation than the originic neumonetheless, well developed glandlike structures are present. In the benign epitheliomia, there is a further step down in respect to differentiation. The basal cell epitheliomias are the least differentiated of the primary equals of the primary equal

of the primary epithelial germ tumors

Bysil cell epitheliomis are not considered carcinomis because they do not metastasize. Then origin from basil cells is doubted because their cells in contrast with basil cells do not possessyntercellular bridges. They are believed to originate from the primary epithelial germ for the following reasons. (1) they occur only in areas where germ for the following reasons (1) they occur only in areas where primary epithelial germ structures (sebaceous glands apoerine glands or harry are found (2) the cells of undifferentiated basal cell epitheliomus resemble the cells of primary epithelial germs as found in embryos from 4 to 6 months old (Fig. 1) (3) structures resembling sebaceous glands and burn are present in many basal cell epitheliomas and (4) basal cell epitheliomas frequently are seen in the same lesion with primary epithelial germ tumors of higher differentiation

TABLE 6—CLASSIFICATION OF THE NEVOID TUNORS OF THE FPIDERMAL APPRIDACES (PRIMARY EPITHELIAL GERM TLMORS)

	WITH SEBACEOUS DIFFERENTIATION	WITH APOCRINE DIFFERENTIATION	With Hair Differentiation
Organic hamar tomas (organic nevi)	Sebaceous nevi 1 Nevus seba ceus (Jadas- sohn) 2 Adenoma se baceum (Prin gle) 3 Senile seba ceous nevus 4 Fordy ce disease	Apocnne nevi	Hair nevi
II Organoid ha martomas (adenomas)	Sebaceous adenoma	Apocrine adeno- mas 1 Syringocysta denoma papilliferum 2 Hidradenoma papilliferum	
II Suborganoid hamartomas (benign epithe 1 omas)	Sebaceous epitheli oma	Apocrine epithelio- mas 1 Syringoma 2 Cylindroma 3 Myo-epithe lioma	Hair epithelio- mas I Tricho-epi- thelioma (epithelioma adenoides cysticum) 2 Calcifying epithelioma
Nonorganic hamattomas Differenti ated basal cell epithel omas b Undifferen trated basa cell epithel omas		Adenoid basal cell epithehoma	heratotic basal cell epithelioma

Some of the terms used to designate groups of tumors require definition

NEVUS This term is used in the literature in two different ways referring either to a tumor composed of nevus cells (nevocellular nevus, pigmented nevus) or to a lesion originating from embryonil cells and composed of mature or nearly mature structures (organic nevi, such as nevus vasculosus, nevus sebaceus, nevus pilosus and nevus verrucosus) In order to avoid confusion, it is advisable to use the term nevus when referring to an organic nevus always with a qualifying adjective, so that nevus without qualifying adjective desig nates a tumor composed of nevus cells According to Jacassolm organic next are tumors and nokhyperplasias because they develop on the basis of an abnormal germ anlage and not by an increase in the size and number of mature structures

NEVOID TUMOR (HAMARTOMA) The term nevoid tumor is used widely as a designation for benign tumors of embryonal origin. How ever, this term lacks conciseness and therefore is unsatisfactory. The term hamattoma appears more satisfactory. This term, derived from the Greek word hamattanein (fail, miss, err) was coined by Albrecht as a designation for tumorlike malformations showing a faulty mix ture of the normal components of the organ in which they occur Van der Valk enlarged the concept of hamartoma to include all be migh tumors of embryonal origin with an organoid structure Since no satisfactory term exists for the entire group of benigh tumors of embryonal origin, the author has suggested that the meaning of the term hamartoma be extended to include all such tumors even those with suborganoid and nonorganic structure. Thus, the tumors aris ing from the primary epithelial germ may be divided into organic hamariomas (organic nevi), organoid hamariomas (adenomas), sub organoid hamartomas (benign epitheliomas) and nonorganic hamar tomas (basal cell epitheliomas)

EPITHELIONA The term epithelioma is used by many authors as a synonym for carcinoma of the epidermis Since, however, the true meaning of the word is tumor of the epithelium the term may be used, as suggested by Jadissohn, as a designation of benign as well as malignant tumors of the epidermis provided that a qualifying as artiguant tumors of the epiderinis provided that a quantificative is added It would perhaps be best if, as Becket his significant, the term optitivelionis be reserved forthering epiderinal tumors and bircinoma fortinalignant epiderinal tumors.

1. TUMORS OF THE SURFACE EPIDERMIS

NEVUS VERRUCOSUS

News vertucous is known also under various other clinical designations, such as hard nevus, epidermal nevus, nevus unius lateralis, linear nevus ketatotic nevus and ichthyosis hystrix



Fig. 133 Nevus verrurosus. There are marked hyper keratosis, acanthosis and papillomatosis. The rete ridges are elongated (X50).

The lessons may be single or multiple and usually are present at birth They consist of serricous growths of brownish color which often show linear combiguration. Large horny excrescences may be present

Histopathology. Nevus vertucous shows hyperkeratosus, papillo matosis and acanthosis with elongation of the rete ridges (Fig. 153). Thus, it has the histologic appearance of a papilloma.

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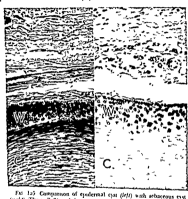
Some of the terms used to designate groups of tumors require definition

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NEVOID TUMOR (HAMARTOMA) The term nevoid tumor is used widely as a designation for benign tumors of embryonal origin. How ever this term lacks conciseness and therefore is unsatisfactory. The term lumintomia appears more satisfactory. This term derived from the Greek word hamartanetn (full miss err) was coined by Albrecht as a designation for tumorlike malformations showing a faulty may ture of the normal components of the organ in which they occur Van der Valk enlarged the concept of hamartoma to include all be nigh tumors of embryonal origin with an organoid structure. Since no satisfactory term exists for the entire group of benign tumors of embryonal origin the author has suggested that the meaning of the term hamattoma be extended to include all such tumois even those with suborganoid and nonorganic structure. Thus, the tumors aris ing from the primary epithelial germ may be divided into organic hamartomas (organic nevi) organoid hamartomas (adenomas) sub organoid hamartomas (benign epitheliomas) and nonorganic hamar tomas (basal cell epitheliomas)

Elthelona The term epithelioma is used by many authors as a synonym for carcinoma of the epidermis. Since however the true mening of the word istumnor of the epithelium the term may be used as suggested by Jadassolin as a designation of benign as well as malignant tumors of the epidermis provided that a qualifying adjective is added It would perhaps be best if as Becker has suggested the term optibelioms be reserved for theman epidermal tumors and barcinoma for malignant epidermal tumors.

cans. These five diseases all show hyperkeratosis and papillomatosis. In typical instances, differentiation is east, but occasionally one is unable to make any more specific diagnosis in these diseases than papilloma. Because the term papilloma is often used in such non



(right) The wall re- of t

amorphous material which has formed by the gradual disintegration of the vacuolated cells (×200)

specific sense in is better to avoid this term as a designation for nexus vertucosus

In typical instances keratosis entilis differe from the pre downward from the form the form as the form the for

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The degree of hyperkeratosis and papillomatosis differs consider ably from lesson to lesson and depends on the sire of the lesson The thickness of the granular layer varies, areas of marked hyperbasis of the granular layer may alternate with areas in which it is attrophed.

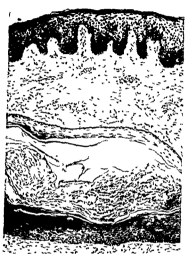


Fig. 154 Epidermal cyst. The wall is composed of true epidermis i.e. squamous granular and horn cells. The cyst is filled with keratin. (×100)

There may be some proliferation of basil cells (as seen in basil cell papilloma) and hyperpigmentation of the basil layer. In rare in stances apocrine gland lumina are found deep in the dermis (see page 346). Nexus cells are absent in pure nexus vertucosus. However, the combination of nexus vertucosus and nexus pigmentosus is not infrequent, and in that case nexus cells are present.

Differential Diagnosis Nevus verrucosus must be differentiated from other types of papillomas namely semile keratosis (keratosis semilis), basal cell papilloma verruca vulgaris and acanthosis nigri Sebackous casts, or steatomas, have a wall composed of epithelial cells that possets no intercellular bridges and do not undergo hera turization Many of the cells are vacuolated because of their transformation into schwerous cells (Fig. 155). The wall never possesses



Fig. 156 Calcined schaceous cyst. The palisading of the basal layer makes it evident that this is a sebaceous cyst. It has ruptured and fibrous issue has proinferated into the lumen (x100).

tete fidges at its periphery. The peripheral cell laver shin + a d a malicide - ~

and cholesterol crystals may be present in the cysts. Calcification oc curs frequently within sebaceous cysts (Fig. 136). As in epidermal cysts a considerable foreign body reaction results when the wall of a sebaceous crist supurers, and the cyst may undergo partial dispure203) Vertuca Vulgaris shows groups of large vacuolited cells in the upper stratum malpighu and the granular layer, and intermitten areas of prirakeratosis (Figs. 118, 119). Acanthosis nigricans, is a rule shows less acanthosis than necus vertucosus and atrophy of the recrudges rather than elongation (Fig. 139).

EPIDERMAL CYST (WEN), SEBACEOUS CYST, MILIUM, DERMOID CYST

Upidermal cysts and sebaceous cysts are often indistinguishable chinically. Both are commonly called wens. On histologic examination, most wens prove to be epidermal rather than sebaceous cysts. In their material, Warvi and Gates found 556 epidermal cysts as comparted with only 3 sebaceous cysts.

Findermal cysts are slow growing, elevated, round, firm, intracutations or subcutaneous tumors, varying from 0.2 to 5 cm in diameter. They occur most commonly on the scalp. As a rule, they are freely movable with the skin. No orifice can be demonstrated in the overlying skin. The material within them is nearly solid and is odorless.

Sebaceous cysts often cannot be differentiated from epidermal cysts on a clinical basis. In general, however, sebaceous cysts are softer than epidermal cysts, and, occasionally, they show a small orifice on their surface. The material within them is semiflucturant and has a ranked odor.

Milia are multiple pithlead sized, whitish, globoid, hard lesions occurring most commonly on the face

Dermoid cysts occur in raie instances in the subcutaneous tissue as soft, round or oval tumors of various sizes. The most frequently observed site is the periorbital region.

Histopathology I PIDERMAI COSTS have a wall composed of true epidermis, i.e., squamous, granular and horn cells (1 ig 154). Rete ridges may be present at the periphery of the wall. The cyst is filled with keratin which frequently is airanged in luminated hyers for calcification are found only in rate instances. When the cyst riptures and the contents of the cyst reiches the dermis, a considerable foreign body grant cell reaction results. The foreign body sextion may cause breaking up and partial disintegration of the epidermal

Malignant degeneration is thre occurring in approximately 15 per cent of cases (Love and Montgomery). If such degeneration occurs, it takes the form of squamous cell carcinoma. Usually, such carcinomas are of low grade malignancy and do not cause metasiass. (Peden)

Selections crists, or steatomas, have a wall composed of epithelial cells that possess no intercellular bridges and do not undergo keraturutum Many of the cells are vacuolated because of their transformation into sebaceous cells (Fig. [55]. The wall never possesses



cro 100 carcined sebsceous cyst The palisading of the bird layer makes it evident that this is a sebaccous cyst. It has ruptured and fibrous tissue has problerated into the lumen (×109)

rete ridges at its periphery. The peripheral cell layer shows a distinct palisade arrangement such as is never seen in epidermal cysts. The cysts are filled with amorphous material which forms by the gradual disintegration of the sebacous cells. Large amounts of cholesterol and cholesterol crystals may be present in the cyst. Calcincianto occurs frequently, within sebacous cysts (Fig. 156). As in epidermal cysts a considerable foreign body reaction results when the wall of a sebacous cyst reputers, and the cyst may undergo partial distinct

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gration. In rare instances, basal-cell epitheliomas have been found arising in sebaceous cysts (Love and Montgomery).

MILIA present a histologic aspect similar to that of epidermal cyst. They are, however, much smaller. Milia are not tumors, like the epidermal and the sebaceous cysts, but are retention cysts caused by the occlusion of a pilosebaceous follicle (Love and Montgomer).

DERMOD CVSTS are lined by an epidermis endowed with rudinen tary sebaceous glands, sweat glands and hair follicles. They conting sebaceous material as well as keratin. In addition, hairs are present in about 30 per cent of cutaneous dermoid cysts. Cartilige and bone are encountered occasionally (New and Eruh).

Differential Diagnosis. For differentiation of epidermal and sebaceous cysts and, particularly, of sebaceous cyst with secondary basal cell epithelioma from calcifung epithelioma, see page 368

KERATOSIS SENILIS

Retatosis sentilis occurs, frequently as multiple lesions, on the fact and the dorsa of the hands in persons past middle life. The lesions

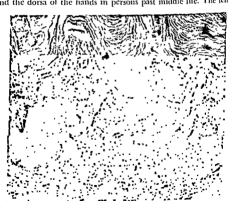


Fig. 157 Keratosis semilis. There are hyperkeratosis and papilloma

usually measure less than 1 cm in diameter and show dry, hard scales firmly adherent to an erythematous base showing hittle or no infiltration Occasionally lesions of sentle kerations show a verticeous surface. In from 20 to 25 per cent of the cases of sentle kerations squamous cell caretinoma develops in one or more of the lesions (Montgomery and Dorffel).

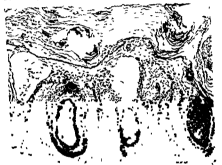


Fig. 158 Keratosis sendis. Clefts and two vesicles are present in the clefts in the clefts.

in the clefts the dermis as (×150)

Cornti cutaneum a variant of keratoris sentlis shows a circum scribed horny excrescence sometimes suggesting the horn of an animal

Histopathology The shortest histologic definition that may be given to keratosis sentlis is that it represents a squamous-cell catci noina Grade 14. It is a precancerous lesson differing only in degree from squamous-cell carcinoma or Bowen's disease.

Hyperketatosis is usually pronounced Mild or moderate papil lomatosis may be present. The stratum malpighii varies in thickness ay alternate. The stratum

of the rete cells. In some of cells, individual cell dyskeratosis, clumping of nuclei and prevalence of mitotic figures are observed, so that the histologic picture approaches that of Bowen's disease (Szodoray) In other cases, the epidermis shows irregular down ward proliferation but without frank invasion (Fig. 157) The his tologic picture then approaches that of squamous cell carcinoma Grade L

Not infrequently, immediately above the basal layer, one observes clefts similar to those seen in Darier's disease (see page 48) These clefts, first described by Freudenthal, may contain a few acantholytic cells In rare instances, even small suprabasal vesicles may be present The basal layer underneath these clefts and vesicles usually consists of cells with large, hyperchromatic nuclei which appear closely set together In addition, this type of atypical basal layer may proliferate into the dermis as short ductlike structures and, furthermore it may surround as cell mantles the upper portion of pilosebaceous follicles and sweat ducts the epithelium of which otherwise appears normal (Fig. 158) (Halter)

The upper dermis usually shows a fairly dense, chronic, inflam

matory infiltrate in which plasma cells are prominent. In instances in which a diagnosis of senile keratosis is made, it often is advisable to section deeper into the block of tissue, because actual progression into squamous cell carcinoma may have taken place in another area

CORNU CUTANEUM is a type of Leratosis senilis with particularly

pronounced hyperkeratosis

vei

tion of cells of the basal cell type (see page 382) (Freudenthal Mont gomery and Dorffel Rutter) For differentiation from squamous cell carcinoma see page 334 from arsenical keratosis see page 155

LEUKOPLAKIA

Leukoplakia represents sentle keratosis of the mucous membranes and occurs on the lips the oral mucosa and the vulva. On the vulva leukoplakia is apt to develop secondary to senile atrophy of the vulsa (see page 162)

Clinically, the lesions consist of white patches which are sharply demarcated, irregular in outline and slightly elevated above the level of the mucous surface. The lesions are often multiple and may be discrete or confluent. In from 20 to 30 per cent of the cases of leukoplakia squamous cell carcinoma supervenes (MacKee and Cipollaro)
Histopathology. In leukoplakia, hyperkeratosis usualli is less pronounced than in keratosis senilis, but the inflammatory infiltrate is

often more intense and may hug the epidermis just as in lichen planus Formation of suprabasal clefts as in sende keratosis does not OCENT

Differential Diagnosis Differentiation of leukoplakia from Jichen planus may be very difficult and occasionally impossible As a rule however the infiltrate in leukoplakia is less severe than in lichen planus and contains a rather large number of plasma cells whereas in lichen planus their number is small. In both diseases the eni dermal-dermal border is apt to have a hazy outline due to the in vasion of the lower epidermis by the inflammatory infiltrate but only leukoplakia shows atypicality of the cells in the stratum malpighii On the buccal mucosa pressure of teeth or dentures may produce pressure calluses which show hyperkeratosis acanthosis and a non specific inflammators infiltrate. They differ from leukoplakia by the absence of atypicality in the epidermis

SQUAMOUS CTI L CARCINOMA (FPIDERMOID CARCINOMA)

Sovamous cell circinoma may occur anywhere on the dam as well ts on the mucous membranes. It may begin as such or develon from t sende keratosis or leukoplakia. Most commonly, the lesion consists of a shallow ulcer surrounded by a nide elevated and indurated border. The ulter often is covered by a crust which conceals a red granular base Occasionally raised fungoid verrucous lesions with ant afteration occur. The latter are usually of a relatively lon grade of malignancy whereas ulcerated lesions may grow rapidly and cause metastases within a short time

Histopathology Squamous-cell cartinoma is a true inter te carri shañ

which the and invade the dermis. The invading tumor masses are composed a ... mous and I

squamous c number of

presses usel

as been carration in the size and the shape of the cells hyperplasia and hyperchromasia of the nuclei absence of prickles keratinization of individual cells prevalence of muotic figures and presence of atypical mitoric figures

Differentiation in squamous-cell carcinomas is in the direction of keratinization keratinization often takes place in the form of horn pearls. The horn pearls are very

of concentric layers of squamous

Legatinization toward the center. The center may or may not show complete keratinization

Compete Aeratinization

Bioders introduced a system of grading squamous cell tatenoma

He established four grades according to the proportion of differentated cells to atypical cells. In Grade I, more than 75 per cent, in

Grade II, more than 50 per cent, in Grade III, more than 25 per cent, and in Grade IV, less than 25 per cent of the cells are differentiated. Since differentiation is in the direction of keraumization. the degree of keratinization is a good guide in grading Broden system of grading is most useful and has been accepted uidely in spite of certain objections raised against it. In the first place, there is of course, a large personal factor in the interpretation of cytologic changes, and, secondly, different degrees of malignancy may be prechanges, and, sectorally, current degrees of mangiane, my be pre-cett in different fields. In regard to the first objection, Broders has suggested to err rather on the side of the higher grade because malig-nant processes (end to be progressive rather thru regressive. In regrid to the second objection, one should examine several sections of every tumor and grade according to the least differentiated portion (I dinundson)

In squamous cell carcinoma, Grade I (Fig. 159), the tumor masses have not penetrated beyond the level of the swert glands. They still show in some areas an intree basal layer at their periphery. In other areas the basal layer has become disorganized and bas disappeared In such ateas the cell masses appert poorly demircated from the sur rounding stroma. The cells of the invading cell misses are predom nantly mature squamous cells with well developed prickles Never theless some of the squamous cells are appical. Horn pearls are present in fairly large number. Some are well developed and have fully keratimized centers others, however show only partral keratimization of their centers and the concentric arrangement of the cells. is not distinct. Besides horn pearls, sheets of partially keratimized cells may be present. The dermis often shows a rather marked in flammatory reaction. It is noteworthy that, in semile kerations and squamous cell caremona Grade I the inflammatory reaction in the dermis is usually much more pronounced than in the more malignant forms of squ timous cell carcinoma. This phenomenon is due to the fact that tissue when invaded by carcinomatous cells is able to defend uself to some extent provided the cells are only modernely malignant, but is occurbelined without any light if the cells are highly malignant (The same observation incidentally can be under also in malignant melanomia and mycosis fungoides—see pages 460 and 485) Metastases do not as a rule occur as long as a squamous-cell carcinoma remains Grade I

In squamous cell carcinoms, Grade II (Fig. 160), the invading cell masses are, as a rule, poorly demarcated from the surrounding stroma. They may invade deeply. Keratimization is much less in evidence than in Grade I. There are only few horn pearls, and those present show, incompletely keratimized centers. A fairly large number of the squamous cells are atypical.



1.0. 159 Squamous-cell carcinoma, Grade J. There is invasion of the derms by epidermal masses. The cells of the invading epidermal masses are predominantly mature squamous cells showing relatively slight asypicality. Several horn pearls are present. The dermis shows a marked inflammatory teaction. (x300).

In squamous cell carcinoma, Grade III (Fig. 161), keratinization is minimal. Horn pearls are not found Instead, keratinization occurs in small cell groups and in individual cells (individual cell keratinization), malignant diskeratiosis see page 499. The majority of cells are atypical. Mitotic figures are conspicuous and often atypical.

In squamous-cell carcinoma, Grade IV (Fig. 162), keratinization is almost completely absent. Nearly all squamous cells are atypical and devoid of prickles. Thus, it is often difficult to arrive at the correct 330

keratinization toward the center. The center may or may not show complete keratinization

Broders introduced a system of grading squamous cell calcinoma He established four grades according to the proportion of different tiated cells to atypical cells. In Grade 1, more than 75 per cent, in Grade II, more than 50 per cent, in Grade III, more than 25 per cent, and in Grade IV, less than 25 per cent of the cells are differ entiated Since differentiation is in the direction of keratinization, the degree of keratinization is a good guide in grading Broders' system of grading is most useful and has been accepted widely in stite of certain objections raised against it. In the first place, there is of course, a large personal factor in the interpretation of cytologic changes, and, secondly, different degrees of malignancy may be pres ent in different fields. In regard to the first objection, Broders has suggested to err rather on the side of the higher grade because malig and processes tend to be progressive rather than regressive In regard to the second objection, one should examine several sections of every tumor and grade according to the least differentiated portion (I dmundson)

In squamous cell carcinoma, Grade I (Fig. 159), the tumor masses have not penetrated beyond the level of the sweat glands. They still show in some areas an intact basal layer at their periphery. In other areas the basal layer has become disorganized and has disappeared In such areas, the cell masses appear poorly demarcated from the sur rounding strong. The cells of the invading cell masses are predominantly mature squamous cells with well developed prickles. Never theless, some of the squamous cells are atypical Horn pearls are present in fairly large number. Some are well developed and have fully keratinized centers, others, however show only partial kera tinization of their centers and the concentric arrangement of the cells is not distinct Besides horn pearls, sheets of partially keratimized cells may be present. The dermis often shows a rather marked in flammatory reaction It is noteworthy that, in senile keratosis and squamous cell carcinoma Grade I the inflammatory reaction in the dermis is usually much more pronounced than in the more malignant forms of squamous cell carcinoma. This phenomenon is due to the fact that tissue, when my ided by carcinomatous cells, is able to defend itself to some extent provided the cells are only moderately defend itself to some extent provided the cells are only moderately malignant, but is overwhelmed without any fight if the cells are highly malignant. (The same observation incidentally, can be made also in malignant melanoma and mycosis fungoides—see pages 460 and 485.) Metastases do not, as a rule occur as long as a squantous cell. carcinoma remains Grade I

In squamous-cell carcinoma, Grade II (Fig. 160), the invading cell passes are, as a rule, poorly demarcated from the surrounding stroma they may invade deeply. Keratuntation is much less in evidence thin in Grade 1. There are only lew horn pearls, and those present show incompletely keratunized centers. A fairly large number of the squa mous cells are atypical.



the 159 Squamous-ell carenoma, Grade 1 There is imassion of the dermis by epidermal masses. The trils of the invading epidermal masses are predominantly mature squamous cells showing relatively slight artipicality. Several horn pearls are present. The dermis shows a marked inflammatory reaction (x/100)

In squamous-cell carcinoma, Grade III (Fig. 161), Leratinization is minimal. Horn pearls are not found. Instead, Leratinization occurs in small cell groups and in indisidual cells ("individual cell kera tinization" "malignini dysk-ratosis," see page 199). The majority of cells are atypical. Mistoite figures are conspicuous and often atypical.

In squamous-cell carcinoma, Grade IV (Fig. 162), keratinization is almost completely absent. Nearly all squamous cells are atypical and devoid of prickles. Thus, it is often difficult to arrive at the correct

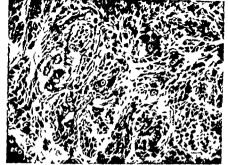


Fig. 160 Squ imous-cell carcinoma. Grade II. The cell masses show much less kerating-tuon than in Grade I. There are only few horn peerly and those present show incompletely keratinged centers https://cells.ore.compicuous (x200).

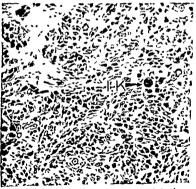


Fig. 161 Squamous-cell carcinoma. Grade III No horn pearls are present. Keratinization occurs only in small cell groups. Many cells are atypical and devoid of prickles. To the right a cell shows. individual cell keratinization. (LK) (x200).

diagnosis as long as individual fields only are studied. The tumor may suggest a malignant melanoma in some cases and a satcoma in others. The latter diagnosis may be pirticularly difficult to rule out when the cells, as occasionally occurs, are spindle shaped ("spindle-cell' squamous cell carexnoma) (Brooks, Underwood, Montgomery and Broders) If, however, sections are subjected to a thorough study, the type of origin from the epidermis and the presence in a few areas

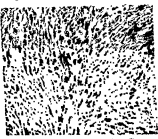


Fig. 162 Squamous-cell carcinoma, Grade IV There is no cudence of kerationation. The epithelial cells appear appical are decord of prickles and elongated so that the tumor suggests a parconna flimost more than a carcinoma. There are numerous/patitotic figures (x200).

of cells showing prickles or partial ketatimization usually establish the diagnosis Squamous-cell cartenoma, Grade IV, is relatively tare in the skin Wany of the reported instances of 'spindle-cell' squamous-cell caremoma occurred in areas of radiodermatitis. In this connection, it may be pointed out that it is not yet established fully whether or not sarcomas can develop following radiodermatis. Most if not all, cases reported as such in the literature tepresent squamous cell cartenomas of the spindle-cell type (Sims and Kirsch, Gentife) (see page 122)

Metastases are rire in squamous-cell carcinoma, Grade I, but common in the other grades. The regional lymph nodes are the first site to be invaded by metastases.

Differential Diagnosis. The diagnosis of squamous-cell carcinoma, although easily made in typical cases, may be difficult at times

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Squamous cell carcinoma must be differentiated from senile kera

Squamous ceil carcinoma must be differentiated from sentie kera tosis, pseudo epitheliomatous hyperplasia and bisal cell epithelioma. The differences between squamous cell carcinoma and sentile kera tosis lie in the degree rather than in the type of changes. In both conditions, one finds atypicality of cells with dyskeratosis of individual cells and downward proliferation of the epidernis. However, in squamous cell carcinoma these changes are more severe, and, in addition, hore pearl formation and actual invasion of the derinis are present No sharp line of separation exists between the two condi-tions, and it is not infrequent to find in a lesion which in general has the appearance of senile keratosis, on serial sections, one or several areas in which the changes have progressed to squamous cell carcinoma.

For differentiation from pseudo epitheliomatous hyperplasia, see below. For differentiation from basal cell epithelioma, see page 380

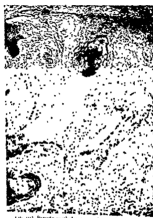
PSEUDO EPITHELIOMATOUS HAPERPLASIA

Considerable thickening and irregular proliferation of the skin, which clinically, as well as histologically, may suggest carcinoma, occurs not infrequently in chronic granulomas, such as bromoderma blastomy costs and granuloma inguinale, and at the edges of chronic ulcers, such as occur after burns, in stasis dermatitis, bisil cell epi thelioma, lupus vulgaris, scrofuloderma, gumma and pyoderma gangrenosum In addition, granular cell myoblastoma is known to Croke quite frequently a pseudo epitheliomitous hyperplisia

Histopathology, Histologically, one observes an epithelial hyper

plasia which may closely resemble squamous cell carcinoma, Grades I or II, and therefore is referred to as pseudo-epitheliomatous hyper plasia. Although squamous cell carcinoma may develop at the edges of chronic ulcers it is likely that some of the cases that have been regarded as such in the past were in reality pseudo epitheliomitous hyperplasia

The histologic picture of pseudo epitheliomatous hyperplasis shows irregular invasion of the dermis by epidermal cell masses with horn pearl formation and often numerous mitotic figures (Fig 163) The penetration may extend even below the level of the sweat glands as penetration may extend even below the level of the sweat glands as isolated fragments of epidermil tissue (Sommerville). However the squamous cells usually are well differentiated, and atypicalities such as individual cell keratinization and nuclear hyperplasia and hyper chromasin are absent. Furthermore in pseudo-epitheliomatous hyperplasia, there often are invasion of the epithelial proliferations by leukocytes and disintegration of some of the epidermal cells, a phe nomenon usually not seen in squamous-cell carcinoma (Winer; Montgomer)) But even when all these criteria are taken into account, it may still be difficult to differentiate between squamous cell carci-



two 103 Freudo-epstheliomatous hyperplasts in bromoderma There is doorn and proliferation of the epidermis analogous to agunnouscell saring. Grade 1 In the field shown it is impossible to rule out tremoma Note, bowever the permeation of the epidermis in many areas by inflammatory cells (X100).

log dıff

noma and .

It is worth remembering to study the influence every section in which one care mona, Grades I or II, i or the granulomatous mis.

.. such evidence is found, one may

Epidermal Tumors

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be dealing with pseudo epitheliomitous hyperplasia rather than with squamous cell carcinomi

"Multipi e, Primary, Sflf Healing Squamous Cell Carcinomata"

In this disorder, which often is familial, there is a continuous ap pearance, especially on the face and the extremities, of papules which grow into nodules, ulcerate and, after a few months, heal with a depressed scar.

Histopathology. The histologic appearance is like that of a well differentiated, keratinizing squamous cell carcinoma, Grade 1 A dense inflammatory infiltrate is present. There may or may not be ulceration, depending on the stage of the lesson.

It is generally agreed that differentiation from squamous cell carci noma is impossible on a histologic basis and depends on the clinical data (Smith, Sommerville and Milne Witten and Zak)

The histogenesis is not clear. However, it is certain that the epi dermal proliferation represents a pseudo-epithehomatous hyper plasia. Whimster assumes that all the downward prolongitions of epithehum can be explained as extreme hyperplasia, probably in firmmatory in origin, of swert ducts and hair follucles.

BOWLN'S DISLAST

Bowen's disease usually manifests itself as a single lesion. It is characterized by a dull red prich of sharp but irregular outline, showing little or no infiltration. Within the prich, there usually are areas of crusting beneath which one finds a granular and oozing surface. The prich slowly spreads by peripheral extension and shows no tendency to healing in its center.

Histopathology. Bowen's disease is an intra epidermal, squimous cell carcinoma, or a squimous cell carcinomi in situ, and not a "pre cancerous dermitosis, under which title it was described originally by Bowen.

The epidermis shows hyperkeratosis with parakeratosis and acan thosis. The rete ridges are elongated and thickened, often to such a degree that the pipillae located between them are reduced to thus strands or are obliterated. However, the bisal layer is intact, and no true invasion can be seen. Throughout the stratum malpighi, the cells lie in complete disorder many are atypical, showing large and hyperchromatic nuclei. Multinucleated epidermil cells containing clusters of muclei are common (Fig. 164). Some cells may show marked vacuolization simulating. Paget cells. However, the intercellular



Fig. 164 Bowen's disease The epidermus is thickened the basal layer is mact. The cells of the stratum malpighi he in complete disorder and many of them are atypical showing large and hyperchromatic mutes. Several multinucleuted cells with clumped mucles and numer to sometimes figures are present in the stratum malpight. (X200)

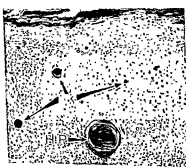


Fig. 165. Bowen's disease. In addition to the changes, as described for Figure 161, there are within the stratum majpights three cells showing individual cell keratinization. (I.K.) and one horn pearl (H.P.) (×200)

338

bridges of these cells are preserved, in contrast with Paget cells, in which they are absent

A common and if present, characteristic finding is the presence of individual cell seratinization in the stratum materials. (Fig. 165)

A common and if present, characteristic finding is the presence of individual cell keratinization in the stratum malpighin (Fig. Such keratinized cells are large and round, have a homogeneous and strongly cosinophilic cytoplasm and a large, irregularly shaped, hyper chromatic nucleus (This individual cell keratinization, which may occur not only in Bowen's disease but also in squamous cell cardinoma, is often referred to as malgiant dyskeratosis in contrast with the benigh dyskeratosis that underlies the formation of corps roads and grains in Dailer's disease) Occasionally, actual horn pearls may occur within the epidermis

The upper dermis usually shows a moderate amount of infimmatory infiltration composed chiefly of lymphocytes and plasma cells

As already stated, in true Bowen's disease the bisal layer is intract. However, in some cases the basal layer ultimately is broken through and a true invasive squamous cell carcinoma results. This may occur at first in only one or a few areas. In order not to miss such areas it is advisable to examine representative sections throughout the entire tissue block. As soon as the invasion of the dermis occurs, the prognosis changes. As long as Bowen's disease remains in the true, intraepidermal stage metastases do not occur. However, when invasion of the dermis has occurred, the likelihood of metastases is rather great. This is due to the fact that if Bowen's disease changes into an in vasive carcinoma, it usually is Grade II or even Grade III with considerable atypicality of the cells and little tendency to keratinization (Kuznitzky and Jacoby)

Differential Diagnosis. In the differential diagnosis senile kera tosis and arsenical keratosis must be considered. Senile keratosis mi) resemble Bowen's disease closely but shows less atypicality of the squimous cells. No sharp line of distinction cun be drawn between the two conditions. Arsenical keratosis may closely resemble either Bowen's disease or senile keratosis but shows as a rule, more vacuolization of the squimous cells than these two diseases.

ERYTHROPLASIA OF QUENRAT

Erythroplasia of Queyrat usually occurs on the glais penis but may be seen occasionally on the prepire the vulva or the oral mucosa. The lesion usually single, consists of a well defined area with a brilliant red, velvety surface and little or no infiltration.

Histopathology. Trythroplasia represents an intre-epithelial squamous cell carcinoma of the mucous membranes and as such is analogous to Bowen's disease of the skin (Pautrier). Progression into

invading squamous-cell carcinoma usually occurs sooner than with Bowens disease of the skin (Sulaberger and Satenstein McDaniel

and Mason Pautner)

Differential Diagnosis Sachs and Sachs as well as Joon recently have described cases in which the clinical appearance was identical with thirt of crybroplasia of Queyrat but in which histologic examination resealed no utalignant changes in the epidermis Instead the derms contained an inflammatory infiltrate composed predominantly of plasma cells. Joon has suggested the diagnostic term balanoposititis chronical circumscripts for these rises Because the two diseases are indistinguishable on clinical grounds it is evident that a diagnosis of crythrophism of Queyrat always requires histologic confirmation.

PAGET'S DISEASE

Pagets disease of the nipple occurs as a rule on and around the nipples of women a few instances of its occurrence on the male breast liste been described (Rubenstein). The lesion is always unlateral Extramanimary Pagets disease which is uncommon occurs on or near the male and the female genitals in the perianal region and in the avillae.

The lesion of Pagets disease consists of a sharply defined slightly infiltrated area of dusk, erythema showing scaling ooning and trius ing. If locited on the breast, the process begins in the impile or the areals of the breast and slowly extends to the surrounding skin. There may or may not be retraction of the inpile.

Histopathology For many years Paget's disease of the mpple was thought to begin in the skin as 1 precamerous lesson that fater be came malignant and then invaded the mammary gland It is now generally accepted that Paget's disease of the nipple is a cancer from the outset and that the mutal lesson is 1 carenoma in situ arising in one or more mammary ducis near their outlets. The primary ductancer extends from the site of origin doinneard to the epithelium luning the actin and upward and outsend to the epithelium luning the actin and upward and outsend to the proferms where it causes the cutaneous lesson. Thus the tumor cells present in the epiderims often referred to as Paget cells are ductal and not epiderimal cells (Pautrier Friser Muir Inglis). At a Tuer stage the cancer breaks through the vall of a duct or actious and infiltrates the connective tissue of the breast.

4 caremonia has been found in the mammary dutts in nearly all cuses of Practs disease and in the mammary glands in most cases. In the few cases where no caremonia was found in the breast the under lying duttal caremonia may have been overlooked because in the

bridges of these cells are preserved, in contrast with Paget cells, in which they are absent

A common and, if present, characteristic finding is the presence of individual cell keratinization in the stratum mulpighii (Fig. 165) Such keratimized cells are large and round, have a homogeneous and strongly cosmophilic cytoplasm and a large, irregularly shaped, hyper chromatic nucleus (This individual cell keratinization, which may occur not only in Bowen's disease but also in squamous cell carci noma, is often referred to as malignant dyskeratosis in contrast with the benign dyskeratosis that underlies the formation of corps roads and grains in Darier's disease) Occasionally, actual horn pearls may occur within the epidermis

The upper dermis usually shows a moderate amount of inflim matory infiltration composed chiefly of lymphocytes and plasma cells

As already stated, in true Bowen's disease the basal layer is intact However, in some cases the basal layer ultimately is broken through and a true invasive squamous cell carcinoma results. This may occur at first in only one or a few areas. In order not to miss such areas it is advisable to examine representative sections throughout the entire tissue block. As soon as the invasion of the dermis occurs the prog nosis changes. As long as Bowen's disease remains in the true, intra epiderinal stage, metastases do not occur. However, when invasion of the dermis has occurred, the likelihood of metastases is rather great This is due to the fact that if Bowen's disease changes into an in vasive carcinoma, it usually is Grade II or even Grade III, with con siderable atypicality of the cells and little tendency to keratinization (Kuznitzky and Jacoby)

Differential Diagnosis. In the differential diagnosis, senile kera tosis and arsenical keratosis must be considered. Senile keratosis may resemble Bowen's disease closely but shows less atypicality of the squamous cells. No sharp line of distinction can be drawn between the two conditions. Arsenical keratosis may closely resemble either Bowen's disease or senile keratosis but shows, as a rule more vacuoli zation of the squamous cells than these two diseases

I RYTHROPLASIA OF QUEYRAT

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Histopathology. Lrythroplasia represents an intra epithelial squamous cell carcinoma of the mucous membranes and as such is analogous to Bowen's disease of the skin (Pautrier) Progression into



Only a



Fig. 168 Pagets disease of the supple Intraductal carcinoma is present in the mammary ducts. The carcinoma is confined within the walls of the ducts (X200)

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ducts the carcinoma may undergo considerable regression with re sultant sclerosis and calcification of the involved ducts (Marx)

Histologic examination of the epideritis reveals in early lesions acanthosis with elongation of the rete ridges, and in older lesions

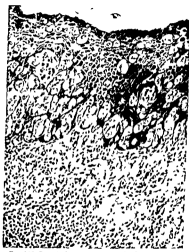


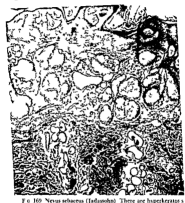
Fig. 166 Paget's disease of the mpple. Low magnification. The epidermis is permetted with numerous Piget cells lying singly and in groups. Note that there is no invasion of the dermis by Paget cells. An inflaminatory influrities is present in the dermis (x200).

thinning and flattening Paget cells are scattered through the epidermis (Fig. 166). They are large, devoid of prickles and at times appear to lie in clear spaces (Fig. 167). Their cytoplasm strains much lighter than that of the adjacent squamous cells. Their nuclei are large, round and pale staining. Paget cells usually are most numerous in the basal layer and may cause disorgranization of this layer. In vasion of the dermis from the epidermis, however, does not occur

2 TUMORS OF THE EPIDERMAL APPENDAGES

NEVUS SEBACEUS (JADASSOHN)

Nevus sebaceus is located most commonly on the scalp or on the face, as a single lesion present since birth. It consists of a circum



and pap llomatos Numerous mature sebaceous glands I e in the upper derm's In the lo er derm's mature apoer ne glands are located (X50)

1 WW!

located (X30) | WWW! scribed slightly raised firm yellos plaque vith a smooth though firrov ed surface Histopathology The tumor is composed of large numbers of ma

or - Frequently do new sebaceous giands The overly ng/epidermis may or - Frequently do n news sebaceus

(R u ep m the dermis bet eath the masses of sebaceous gland lobules (Fig 169) The pres ence of abortive hair follicles has also been noted on several occasions In some cases, Paget cells are so numerous throughout the epiderims that the normal squamous cells show signs of injury. They are compressed and deformed and may form only a network, the meshes of which are filled with Paget cells (Fig. 166)

The dermis shows in Paget's disease a moderately severe chronic

inflammatory reaction

Histologic examination of the mammary ducts nearly always shows malignant changes in some of them. At first, the carcinoma is intraductal and the tumor cells are confined within the walls of the ducts (so called comedo carcinoma) (Fig. 168). Ultimately, the tumor cells invade the connective tissue. From then on, lymphatic spread and metastases occur.

FARMAMMAN PAGETS DISFASE, as a rule, presents the same epidermal changes as Paget's disease of the nipple, although in lesions located in the perianal region the Paget cells often contain a considerable amount of mucin. In most reported cases, an underlying carcinoma of apocrine ducts and glands has been found (Parsons, Foraker and Millet, Zoon and Gelpke). This is in accordance with the fact that the mammary gland is a modified apocrine gland. The presence of mucin within the Paget cells of some of the cases located in the perianal region may be the result of mucous metaplasia of apocrine glands, but Whinster suggests that the primary cancer may be one of ectopic mucous glands rather than of apocrine glands.

Differential Diagnosis. Paget's disease of the nipple must be dif ferentiated from Bowen's disease. In Bowen's disease, large, vacuolated epidermal cells may also occur, but, in contrast with the Paget cells, they often possess prickles Furthermore, one observes clumping of nuclei within multinucleated epithelial giant cells and individual cell keratimization in Bowen's disease but never in Paget's disease. In cases in which the Paget cells are concentrated in the lower epidermis, the resemblance to an amelanotic junction nevus or an early malignant melanoma may be very great because in both diseases the characteristic cell is large and vacuolated (Stout) Allen believes that many cases of Paget's disease and especially of extra mammary Paget's disease are wrongly thus diagnosed and in reality are junction ness. The most important points of differentiation are first, the absence of invasion of the dermis by the tumor cells in Paget's disease, and second, the presence of minute amounts of mela nin in at least some tumor cells even in apparently amelanotic junc tion nevi and malignant melanomas. The melanin is best demon strated by the use of a silver stain Carrying out of the dopa stain (see page 12) also aids in the differentiation

2 TUMORS OF THE EPIDERMAL APPENDAGES

NEVUS SEBACEUS (1/DASSOH/)

Nexus sebaceus is located most commonly on the scalp or on the face as a single lesion present since birth It consists of a circum



Fig. 169 Nevus sebaceus (ladassohn). There are hyperkeratos s. and papillomatous Numerous mature schaceous glands I e in the upper dermis. In the lower dermis mature apoetine glands are located (×50) 1 well

scribed slightly raised firm yellow plaque with a smooth though furroy ed surface Histopathology The tumor

ture or nearly mature sebaceous or may not show hyperkeratosis crine glands have been describe a occurring in nevus sebaceus (Robinson koch Pantrier) They are located deep in the dermis beneath the masses of sebaceous gland lobules (Fig. 169). The pres ence of abortive fair follicles has also been noted on several occasions

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(Ostrowski). Koch and Pautrier believe that the frequent presence of apocrine glands is evidence that nevus sebaceus develops from primary epithelial germs (For a discussion of the primary epithelial germ, see pages 4 and 318.)

The presence of a basal cell epithelioma within a nevus sebaceus is not uncommon (Fig. 170) (Ostrowski; Robinson; Szodoray, Pautrier,



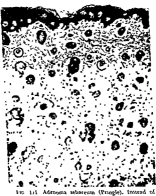
Fig. 170 Nevus sebaceus (Jadassohn) within which a basal cell epithelioma has arisen. The nevus sebaceus is on the right side, the basal cell epithelioma on the left. (×25)

Savatard) In rare instances, a squamous-cell carcinoma may develop from the epidermis overlying the nevus sebaceus (Parkin).

ADENOMA SIBACEUM (PRINGLI) (TUBEROUS SCLEROSIS)

Adenoma sebaceum (Pringle) usually represents only part of a widely dissemunated tissue malformation which results in the formation of tumors in vota ecodernic and mesodernic structures. Tumors are ound, aside from the skin, notably in the brain, the retina, the bidney and the heart. The tumors in theoram, which are referred to as tuberous selerous, are gluomas. Those of the fetina have the same histologic appearance as the brain tumors (Kreim). The tumors of the heart usually are rhaddomyomas (Pratt-Thomas). Those of the Vidney may be angiomas, fibromas, adenomas or mixed tumors; they may become malignant (Butterworth and Wilson).

The cutaneous lessons of adenoma schaceum (Pringle) occur on the fare_particularly in the <u>malar resum</u>. They consist of numerous small papules and modules which are yellowish brown in color and frequently show telanguectives on their surface.



tin 1:1 Adenoma sebateum (Pringle), Instead of showing excessive development of sebaceous glands, as is usually the case, this lesion shows a large number of immature hair structures (X50)

Histopathology. The term adetooms e-baceum is a mistomer. Since the sebaceous structures are not adenomators but fully or almost fully developed and abnormal merely by their presence in excessive number, the lesion represents an organic nesus, rather than an adenoma (see Table 6 - come 2006.)

In addition to a

found Usually, those hair structures are somewhat immature (Fig 171) (Butterworth and Wilson, Good and Garb)

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(Ostrowski). Koch and Pautrier believe that the frequent presence of apocrine glands is evidence that nevus sebaceus develops from primary epithelial germs. (For a discussion of the primary epithelial germ, see pages 4 and 318.)

The presence of a basal-cell epithelioma within a nevus sebaceus is not uncommon (Fig. 170) (Ostrowski; Robinson; Szodoray, Pautrier;

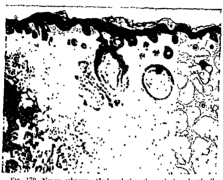


Fig. 170. Nevus sebaceus (Jadassohn) within which a bisil cell epithelioma has arisen. The nevus sebaceus is on the right side, the basal cell epithelioma on the left. ($\times 25$)

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Adenoma sebaceum (Pringle) usually represents only part of a widely disseminated tissue malformation which results in the formation of tumors in both ectodermic and mesodermic structures. Trumors are dound, aside from the skip, notably in the brain, the retina, the kidney and the heart. The tumors in the brain, which are referred to as tuberous selerosis, are gliomas. Those of the tetina have the same histologic appearance as the brain tumors (Kveim). The numors of the heart usually are rhabdomyomas (Pratt-Thomas). Those of the didney may be angiomas, fibromas, adenomas or mixed tumors; they may become malignant (Butterworth and Wilson).

SEBACEOUS ADENOMA

An adenoma may be defined as an organoid tumor consisting of circumscribed proliferations of incompletely differentiated glandular structures. If this definition is adhered to, sebaceous adenoma is a very rare tumor. Many tumors described in the literature as sebaceous



110. 110. senacous adenoma i he tumor is composed of lobules which are irregular in size and shape. Two types of cells compose the lobules, generative and sebaceous cells (X100)

adenoma are in reality sebaceous nevi. Examples of sebaceous ade noma have been described by Renmann, Pautrier, Woolhandler and Becker, and Lever, among others

Sebaceous adenoma occurs as a smooth, firm, round, elevated, often slightly pedunculated, tumor. In most reported cases, the lesion was solitary, located on the free or the scalp, and measured less than 1 cm in diameter.

Histopathology. On histologic examination, the tumor is sharply demirrated from the surrounding tissue and is usually surrounded by a connective tissue capital? It is composed of broules which are triegular in sure and shipe (Fig. 172). Two types of cells are present.

SENILE SEBACTOUS NEVUS (SENILE SEBACEOUS HYPLRPLASIA

Semile sebaceous nevus occurs on the face, chiefly on the forehead, in persons past middle life, and consists of either one or a few small vellowish, translucent nodules

Histopathology. This lesion, like nevus sebaceus (Jadassohn) and adenoma sebaceum (Pringle), is composed of large numbers of ma ture or nearly mature schaceous glands. Differentiation of senile sebaceous nevus from the other two lesions on histologic grounds is therefore, often impossible In contrast with nevus sebaceus and Adenoma sebaceum (Pringle), however Apportine glands or immature Mair follicles are never found

Senile sebaceous nevus is regarded by some writers as a senile hy perplasia of sebaceous glands (Gilman, Woolhandler and Becker), while others consider it as a delayed, sende organic nevus or himar toma (Gans) The cucumscribed nature of the lesions makes the latter view more likely than the former

FORDYCE'S DISFASE

In this condition, groups of minute, yellowish, globoid lesions are observed on the vermilion border of the lips or on the oral mucosa

Histopathology. Fordyce's disease has as pathologic substrate the presence of schreeous glands in areas where they are normally absent It thus represents a sebucious nevus or hampronn

Histologic examination reveals groups of mature schreeous lobules located in the upper dermis Some lobules lie free in the dermis others he at the end of downward proliferations of the overlying epiderinis, and still others are connected with the epiderinis by true sebucous ducts (Chambers). However chairs are never found

APOCRINE GLAND NEVUS

Organic nevi composed only of apocrine glands do not occur Aportine gland structures however, frequently are present in nexus sebaceus (Jidissohn) (see page 343) and occasionilly in nexus ver rucosus (sce page 322)

HAIR, NEVUS

Furely hair next occur, but hore commonly they appear in conjunction with other next old lesions, for instance with nexts sebrecus (Jadassohn), nexts vertucosits and nexts pigmentosits. Some cases of adenoma sebaceum (Pringle) are hair next rather than sebaceum sext (see page 345)

schaceous lobules. In most lobules, however, the two types of cells occur in approximately equal proportions, often arranged in such a way that groups of schaceous cells are surrounded by generative cells Larger lobules may contain in their center cystic spaces formed by the decomposition of mature sebaceous cells

Wat stains reveal the presence of fat in the sebaceous and transi

tional cells and in the cystic spaces

SYRINGOCYSTADENOMA PAPILLIFERUM (NEVUS SYRINGOCYSTADENOMATOSUS PAPILLIFERUS)

This tumor represents an apocrine gland adenoma with differ egitation predominantly toward apocrine dista.in contrast with Midradenoma papilliferium in which differentiation is directed predominantly toward apocrine glands



Fig. 174 Syringocystadenoma papilliterum. High magnif ation of Figure 173. The vills are lined by two rows of cells. The outer row is composed of <u>small</u> cubo dat cells (mose-publical cells) the inner row of Ji gh cylindrical cells (secretory cells of aportine glands) (2200)

Clinically, syringocystadenoma papilliferum occurs no all a single terr on all tered th

cysts can

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in the lobules. The cells of the first type are identical with the cells present at the periphers of normal sebaceous glands and resemble the cells of which basal cell epithelionars are composed. They have

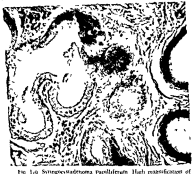


Fig. 173 Syringocystadenoma papilliferum Low mag miferition. A cystic invagination extends down and from the epidermis. Numerous villuslike projections extend into the lumen of the cystic invagination. A group of apoetine glands (14) is present in the left lost eromer (x50).

been orlled generative cells by Grynfeltt (see page 17) The cells of the second type are mature sebucous cells. They have developed from the generative cells. In addition there are cells an a transitional stage of differentiation. The distribution of the generative and the sebaceous cells within the lobules varies. Some lobules contain predominantly generative cells and thus resemble basic cell epitheloma. Other lobules contain mainly sebaceous cells and resemble mature

ence of active secretion. These cells represent secretory cells. Cel

Meneath the cystic invaginations deep in the dethils one finds tougs of glandular himens (Fig. 175). Their secretory activity (de apuation secretion) clerify labels them as apocrine glands (Fig. 176) (Tappenner Appel Grund) Connections of the apocrine glands.



the aportine glinds in Figure 170. The secretory cells of the aportine glinds in Figure 170. The secretory cells of the aportine glinds show evidence of active secretion (decapitation secretion.) (×400)

with the cystic imaginations in the upper dermis can be traced with out difficulty.

In most cases a dense inflammatory infiltrate is present in the

in most cases a section quantitative present in the upper definite and especially in the villi. This infiltrate contains a large percentage of plasma cells and in some cases is composed almost exclusively of plasma cells.

The association of syringocystadenoma papilliferum with nevus sebacus (Dofffel Marcus and Wooldridge Grund) or with hasal cell epithelioma (Dofffel Reuterwall) has been noted on several occasions

Differential Diagnosis For differentiation from Darier's disease in which villi are also sometimes very prominent, see page 50

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Histopathology. The Epidermis shows acanthosis and papillomatosis. Cystic invaginations extend downward from the epidermis (Fig 173). These invaginations represent greatly dilated apportine ducts



Etc. 175 Sytungocyusukenoma papulliferum Los, mogunustion In the upper dermis there are several gistic intagnations filled with villuslike projections. A mixedj inflamatory in filtrate containing many plisma cells is present around the cystic intagnations. The lower dermis contains numerous apoerine glands (x50).

Numerous villuslike projections extend into the lumina of the in vaginations. The cystic invaginations, as well as the villi, are fined by the rows of cells (Fig. 174). The outer row is composed of small cuboidal cells with deeply staining nuclei. These cells are immature myo epithelial cells. The inner row is composed of lugh, cylindrical cells which have large, oral, pale-staining nuclei and may show evi





Gates Warren and Warvi have pointed out that hidradenoma papilliferum closely resembles the papillary cystadenomas of the mamming gland It should be remembered that, phylogenetically, the mamming gland is an apocrine gland

SEBACEOUS EPITHELIOMA

Clinically the tumor has no characteristic appearance. Usually, it presents itself as a <u>solitary</u> small nodule or plaque. It is an uncommon type of tumor. Examples have been described by Grynfelit, Biberstein Milian, Perin and Brunel, and Lever.

Histopathology. In degree of differentiation sebaceous epitheli oma stands between sebaceous adenoma in which there are typical sebaceous lobules, and cystic basal-cell epithelioma, in which there

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HIDRADENOMA PAPILITERIM

This tumor occurs almost exclusively on the labia majora and on the perineum of women as a solitary intracutaneous lesion covered by normal epidermis. It usually measures only a few millimeters in diameter Malignant degeneration does not occur



Fig. 177 Hidradenoma papilliferum Low magnifica t on The ti mor cons sts of a large cistlike lumen into h ch numerous interlacing villi project (×50)

Histopathology The tumor represents an adenoma of apocrane glands (Anderson Winer) It is located in the dermis shows no con nection with the epidermis and is well encapsulated. It is composed of a large cystlike lumen into which numerous interlacing villi pro ject (Fig 177) The will of the cyst as well as the villi are lined usually by a single layer of high cylindrical cells. These cells have a

differentiation is directed mainly loward apocrine duct cells, differentiation in solutionary is directed mainly toward apocrine, gland cells and many o epithelionar mainly toward apocrine may epithelial

yellowish nodules, the size

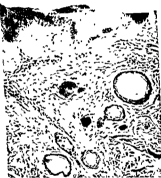


Fig. 180 Syringoma. The deright contains several small cysic ducts: The walls of most ducts are lined by <u>two zone</u> of <u>equilical zeils</u> <u>Colloudal</u> maternal falls the <u>lumina</u>. Two of the ducts have commanded to the ducts have commanded to the contains after the property (X200).

of a punisged are found around the eyelds on the chest, on the abdomen and on the anterior aspects of the thighs, but occasionally also elsewhere on the skin

Histopathology The derinis contains numerous, small, cystic ducts (Fig. 180). The walls of the ducty are lined usually by two rous of epithelial cells in most instruces these cells are flat and are it compressed.

evidence of act

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is but little differentiation toward sebaceous cells (See Table 6 page 319) As Loos has put it sebaceous epithelioma giows like a basal cell epithelioma but its cells have undergone considerable differentiation toward sebaceous cells

The tumor is composed of irregularly shaped tell masses which are scattered through the upper decimis (Fig. 179). The majority of cells



Fig. 179 Sebreous epitheliama. The tumor is composed of irregularly shaped cell masses. The majority of cells are of the same type is in bisalcell epitheliama but many cells (SC) show differentiation to and sebaceous cells (X100).

are of the same type as the gells in basal cell epitheliona. However a fairly large number are stansinoral cells showing beginning fatts transformation of their cytoplasm. Groups of faithre sebaceous cells in the centers of the cell masses. Cysts formed by the disintegration of sebaceous cells may or may not be present in the center of some of the tumor masses.

SURINGOMA

Three types of apocrane epitheliomas occur syringonia cylin droma and myo epithelioma (see Table 6) While in syringonia Besancon Riehl Homma and Fscher Wendlberger) (2) its simulta neous occurrence with rudimentary hair structures or with tricho epithehoma a tumor of hair structures (Fischer Lever) (3) its ap pearance at puberty to en apocrine glands first begin to function (kyrle Wendlberger) and (4) the prevalence of the lessons in areas of the skin where apoerine glands occur or used to occur phylogeneti cally (Wendlberger)

CYLINDROMA

This disorder which is often hereditary is characterized by the presence of numerous reunded smooth tumors of various sizes on



aport ne gland cells (X900)

the scalp Occasionally a few tumors are present also on the facand the upper trunk. The lesions begin to appear in early adulthood and increase in number and size throughout life. They vary in size from a few millimeters to several centimeters and by their arrange ment in groups resemble bunches of grapes or tomatoes. The tumor man and Besancon; Riehl; Homma and Escher; Wendlberger). The lumina of the ducts are filled with a colloidal material Some of the cystic ducts possess small, comma-like tails of epithelial cells, giving them the appearance of tadpoles In addition, there are solid strands of epithelial cells independent of the ducts. The cells composing the

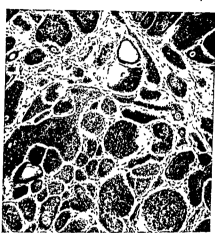


Fig. 181 Cylindroma, Low magnification. The tumor is composed of various sized islands of epithelial cells. The islands are sur lower quadrunt contain droplets of hyalin (×75) rounded by a hyaline membrane. Several of the islands in the left

solid strands have the same appearance as the cells in basal cell epithelioma. Rudimentary hair structures are found occasionally in the lesions (Fischer).

Several authors have reported the simultaneous occurrence of syringoma and tricho epithehoma (epithehoma adenoides cysticum) in the same patient (McDonagh; Weidman and Besancon) and even in the same lesion (Ingels; Lever).

The reasons given by various authors for the apocrine genesis of syringoma are: (1) the presence of active secretion (Weidman and

Besancon Richl Homma and Escher Wendlberger) (2) its simultaneous occurrence with rudimentary hair structures or with trichoeputhelioma a tumor of hair structures (Fischer Lever) (3) its appearance at <u>puberty</u> when apocrine glands first begin to function (kytle Wendlberger) and (4) the prevalence of the lesions in areas of the skin where apocrine glands occur or used to occur phylogenetic rally (Wendlberger)

CYLINDROMA

This disorder which is often hereditary is characterized by the presence of numerous rounded smooth tumors of various sizes on



Fig. 18° Cylindroma. H gh magn ficat on of F e re 181 'r

appearing gland cells ($\times^{0}00$)

the scaip Occasionally a lew tumors are present also on the Jace and the upper-trunt. The lesions begin to appear in early adulthood and increase in number and sire throughout life. They vary in size from a few millimeters to several tenumeters and by their arrange ment in groups resemble bunches of grapes or tomatoes. The tumors

man and Besmoon, Riehl, Homma and Escher, Wendlberger) The lumina of the ducts are filled with a colloidal material. Some of the cystic ducts possess small, comma like tails of epithelial cells, giving them the appearance of tadpoles. In addition, there are solid strands of epithelial cells independent of the ducts. The cells composing the

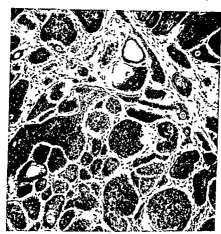


Fig. 181 Cylindroma I ow magnification. The tumor is composed of virious sized islands of epithelial cells. The islands are sur-lower quadrant contain droplets of hyalin. (x75), rounded by a hyaline membrane. Several of the islands in the left

solid strands have the same appearance as the cells in basal cell epithelioma. Rudimentary hair structures are found occasionally in the lesions (Fischer).

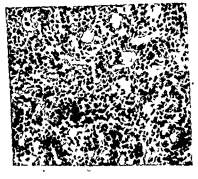
Several authors have reported the simultaneous occurrence of syringom; and tricho epithelioma (epithelioma adenoides cysticum) in the same patient (McDonagh, Weidman and Besancon) and even in the same lesion (Ingels, Lever)

The reasons given by various authors for the apoctine genesis of syringoma are (1) the presence of active secretion (Weidman and

Myo-epithelioma (Myo epithelial Sweat Gland Tumor) nd (3) its simultaneous occurrence with tricho epitheliotia a tumor of hair structures

MO EPITHELIOMA (MO FPITHELIAL SWEAT GLAND

Uso epithelioma occurs is a rule as alsoluary tumod occasionally however several lesions are present. The tumors present themselves



as firm introdutineous nodules and usually measure between 0.5 and 2 cm in diameter although they may be larger. The overlying skin is either normal or shows slight erithema. Myo-epithelioma is a rather uncommon tumor Examples have I een described by Sheldon Balog, Harry and Lever

Histopathology Histologic examination reveals a sharply circum scribed usually encupsulated essentially solid tumor. The epithelial cells of the tumor are arranged usually in intertuining bands but orcisionally in lobular masses. The same two types of cells are pres ent as in cylindroma namely secretory and myo-epithelial cells but

may cover the entire scalp like a turbin. For this reason, they are referred to occasionally as turbin tumors.

Histopathology. The tumors are composed of numerous, variously sized islands of epithelial cells. These islands are surrounded by a highline membrane. In addition, droplets of hyalm are present in some of the islands (fig. 181). You types of cells compose the islands (Stillrans, Savatard, 7akon, Lever). Cells with large, oxal, light striping nucles he in the centers of the islands, while cells with small, round, tark strining nuclei are present in palisade arrangement at the pertiplery of the islands and also scattered between the cells of the first type (fig. 182). The cells of the first type may be regarded as cells differentiating toward secretory cells, those of the second type as cells differentiating toward myo epithelial cells of apocrine glands. The secretory cells always outnumber the myo epithelial cells. The dividual which is present around and within the islands appears to be produced by the myo epithelial cells, since it is always found in apposition to them. It is particularly well demonstrated by straining with methylene blue.

In some cases, a thorough inspection of the histologic sections will reveal the presence of a femediandular lumina lined by two layers of cells an inner actively secreting layer and an outer myo epithelial layer (Stillians, Lever) (Fig. 182). These glandular structures strongly

resemble apoerme glands

Several authors have observed connections of the immor misses with hirr folkeles (Watanabe, Stillians Schlammadinger). The simultaneous presence of cylindromi and tricho epitheliomi (pitheliomi adenoides cysticum) in the sime pittent has been observed repeatedly (Watanabe. Schlammadinger. Schuermann and Weber, Smatrid). Cylindromas neithy alarys remain benign. Two cases, however, are on record in which indignant degeneration of the immors occurred with metistases to the lymph nodes in one case (Liusecker) and to internal organs as well in the other (Luger). The areas of anthomatic degeneration showed in the cell lobuler polymorphism of the mater numerous/mitotic figures. loss of the hyrline sheath, loss of pilisading at the pyriphery and im asson into the surrounding stromi.

The distogenesis of cylindromi is not yet clearly established. Some

The distogenesis of cylindrom's not yet clearly established Some authors (Coenen, Davies) regard it as a succi gland tumor, others (Watapabe, Sullians, Schlammrdunger) as a hair folloide tumor and still others (Balog Schuermann and Weber) as an apactine tumor In favor of the apactine genesis of cylindrom are (1) the presence of two types of cells namely, secretory and myo epithelial cells (2) the occasional presence of actively secreting glandular lumina

and (3) its simultaneous occurrence with tricho-epithelioma, a tumor of hair structures

MIO EPITHELIOVIA (MIO EPITHELIAI SWEAT GLAND TUMOR)

Myo epithelionia occurs as a rule as alsoluare tumod occasionally, however several lesions are present. The tumors present themselves

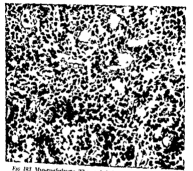


Fig. 183 Myo-epitheliona. The epithelial cells are stranged in intertioning binds. Two types of cells can be seem-secretary cells lie in the center of the bands and trad to be arranged around lumina. Myoepithelial cells he at the prophery of the binds (x400).

ic Frm

Histopathology Histologic examination reveals a sharply circum scribed usually encapsulated essentially toled tumor. The epithelial cells of the tumor are arranged usually in intertwining bands but occasionally in lobular masser. The same two types of cells are present us in cylindroma namely secretory and myo embelled cells.

in contrast with cylindroma, the majority of cells are myo epithelial cells. The secretory cells lie in groups and frequently are arranged around lumina which are usually small.

In myo epitheliomas with bandlike arrangement of the epithelial cells, the secretory cells are located in the center and the myo epi

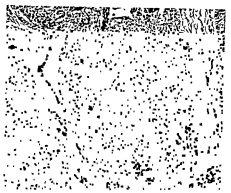


Fig. 184. Myo-epithelioma. The epithelirl cells are arranged in lobules. The lobule shown is composed largely of myo epithelirl cells but lumina lined with secretory cells are scattered through the lobule (x200).

thehal cells at the periphery of either tubules or solid binds (Lig 183). The myo epithelial cells may be seen prohiferating irregularly into the stroma. In myo epitheliomas with lobular arrangement, the lobules are composed largely of myo epithelial cells, but throughout the lobules one finds luming lined with secretory cells (Fig. 181).

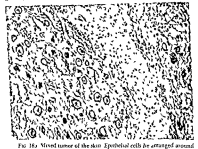
Some of the tumors possess a considerable amount of stroms of hadine appearance and thus resemble the so called mixed tumors of the salivary gland type

MINED TUMOR OF THE SKIN

Mixed tumors of the salivary gland type occasionally have been described as occurring in the skin (Hirsch, Highman, Morehead,

Lennox Pearse and Richards) Their Einical appearance is the same

as that of man couthelioma Hutopathology Mixed tumors of the skin show epithelial cells arranged in nests and strands as well as around small lumina (Fig 185) Either one or two layers of epithelial cells are present wound the lumina. In addition epithelial cells may be seen proliferating stregularly from the luming into the stroma The majority of epi



small tubular lumina and scattered through the stroma. The stroma shows mucoid and byaline degeneration (×100)

thehal cells are spindle shaped and resemble myo-epithelial cells (Moreherd) Those located around luming have the appearance of clandular cells The stromy usually is abundant and shows hyaline and muroid degeneration

Some authors regard the mixed tumors of the skin as of sweat Lind origin (Hirsch Lennox Pearse and Richards) It is the author's belief that most if not all mixed tumors of the skin represent myo epitheliomas. They differ from min epitheliomis only by showing a greater amount of stroms and more marked degenerative changes in the stroma.

GLEAR CELL. MAG EPITHELIOMA

The clinical appearance of this tumor is that of a solutary nodula usually covered by intact skin but occasionally discharging serous material

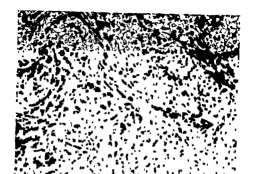


Fig. 186 Clear cell myo epithelioma. The tumor is composed of two types of cells Justform myo epithelial cells and suboidal clear cells." (x200)

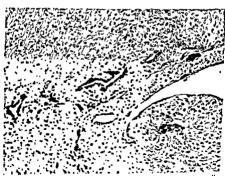


Fig 187 Clear-cell myo epithelioma. There are numerous clear cells and in addition, tubular luming lined by a single layer of secretory cells. (X200)

Histopathology. The tumor has a lobular structure. Two types of epithelial cells cun be recognized dissiform mio-epithelial cells with deeply bisophilic nuclei and clear cells which are cuboudal have a distinct cellular membrane very clear cytoplasm and a round nucleus (Fig. 186). The clear cells seem to develop from the myo epithelial cells (Lever and Cisileman). Varying amounts of glicogen any present in the clear cells. In some timors there are groups of veraitinged cells cysts and glandular structures. The glandular structures are rubular and their lining cells often show. decapitation

secretion as observed in aportine glands (Fig. 187).
While Lever and Castleman regard this type of tumor as a variant of myo epitheliona. Liu because of the presence of glycogen thought that it was derived from the outer bars sheath which normally concums abundant amounts of glycogen. However, glycogen is found so commonly in joung cells and in tumors that its presence crimto be considered as a reliable criterion for the histogenetic classification of neoplasms.

TRICHO EPITHELIOMA (FPITHELIOMA ADENOIDES CISTICUM MULTIPLE BENIGN CISTIC FPITHELIOMA)

The name tricho epithelioma is prefetable to the other names listed above because it indicates that differentiation in this tumor is directed toward four structures

The disorder begins as a rule at the age of pubert, and is frequently heredity. It is characterized by the presence of numerous punhead to pea sured rounded vellowsh or pinh nodules on the face and occasionally on the upper-trunt. A few telanguectatic vessels are often present on the surface of the larger lesions. Occasionally one or several lesions become ulcerated because of change into a basal-cell epithelioma.

Histopathology On histologic examination tricho-epithelionia appears as a well-accumorabled immore/from cysts represent the characteristic lesion. They consist of a fulls, ferginning leners sur-rounded by a shell of finitened. Justif cells without/prickles (Fig. 188). The keratinnianon in the form cysts is althout-and compilere noise radiual and incomplete as in the horn pearls of squ'immousceil exteriorism. This process corresponds to the abrupt development of the horn cells of the hur from the hur matrix cells (which are also cells without prickles). It may be concluded therefore that the cells surrounding the horn civis are lain matrix cells and that the horn custs represent attempts at base shift formation (Lever). Occasionally, one sees prickle cells around some of the horn cysts. They represent outer hair sheath develops from

prickle cells at a time when the hair germ has already advanced to a rather high stage of differentiation the presence of prickle cell around the horn cysts is evidence of rather high differentiation

In addition to horn cysts Yrregularly shaped islands and intertwining strands composed of basal cells are present Such areas are indistinguishable from basal-cell epithelioma (Goldman) Abortito.

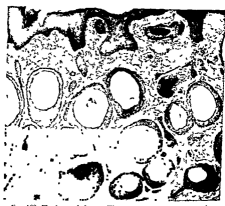
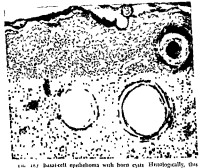


Fig. 188 Tricho epithelioma. The tumor contains numerous homers in the center near the epidermis two rudimentary hairs can be seen (×100)

hair papillae and hair shafts are seen occasionally. Since hair papillae contain a high concentration of alkaline phosphatase (Hardy) their presence can be well demofistrated by the use of the Gomory stam for alkaline phosphatase Calcification of the horn cysts may occur and evoke asforeign body grant cell reaction in the adjacent connective tissue.

Tumors histologically in any intermediate stage between tricho epithelionia and basal cell epithelionia occur (Summerill and Hut ton Trienkle). As just stated a typical tricho epithelionia is char accerized by cricumscribed growth and by the presence of many hom cysts and of basal cell masses however basal cell epithelionia may

also show horn cysts (Fig. 189) (see page 379) and may be circum scribed in growth. Since thus no sharp line of demarcation exists between tricho-epithelioma and bisal-cell epithelioma histologically, it may be necessary, in order to arrive at a diagnosis in a given case, to have knowledge of clinical data, such as the number and the distribution of the lesions and the age at which the lesions first ap



110. Dasai-cell epithenoma with hord cysts 11stologically, this tumor is in an intermediate stage of differentiation between basal cell epithelionia and tricho-epithelioma. Clinically the lesion was a basal cell epithelioma. (X200)

peared. The close relation between the two types of tumors is attested further by reports of caset in which one or several of the lestons of trutho-epithelioma after having persisted as such for many years, developed into ulcers with the histologic picture of basal cell epithelioma (Mamson Little, Saviatard). This close relationship of the two types of tumors can be explained best by assuming that they have seminate general regions (Level) and that they differ only in the degree of maturity of their cells (Adamson). Since cells of various maturity may occur in the same lesion, one may find in tricho-epithelioma areas consistent with the histologic picture of basal-cell epitheliom and vice tests, also, if active growth occurs in a lesion of tricho-epithelioma, the newly formed cells may

be more embryonal than the older cells and the lesson may grow as a basal cell epithelioma

A close relation exists not only between tricho epithelioma and basal cell epithelioma, but also between tricho epithelioma and other types of benign epitheliomas such as syringoma and cylindroma Tricho epithelioma may occur with syringoma or cylindroma in the same patient (see pages 356, 358)

CALCIFYING EPITHELIOMA (MALHERBL)

Calcifying epithelioma is a solutary hard, deep seated tumor which is covered by normal skin. It occurs most frequently on the face and



Fig. 190 Calcifying epithelioma (Matherbe) The tumor consists of lobules embedded in a connectine tissue stroma Tho types of cells compose the lobules—basophilic cells—and shadow cells. The basophilic cells resemble the cells of basal cell epithelioma. The shadow cells show a central unstained shadow at the site of the muleus in the center of the field one cap see transformation of the basophilic cells into shadow cells. The stroma contains numerous foreign body, giant cells (X100)

the upper extremities. The size is from 0.5 to 3 or even 5 cm in diameter.

Histopathology Calcifying epithelioma is a sharply demarcated, often encapsulated tumor and usually located in the lower dermis

or in the subcutaneous fat. The histologic appearance is character istic. Embedded in a connective tissue stroma <u>variously shaped</u> masses of epithelial cells are present. As a rule, that tipes of cells—basophilic cells and shadon cells—compose theseymasses (Fig. 1900) Occasionally basophilic cells are absent. The Vasophilic cells greatly resemble the cells of basal cell epithelioma. They possess

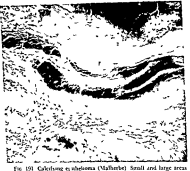


Fig. 191 Calcilying et inhelionia (Mainerbe) Small and large areas of calcification are present within the lobules of shadow cells (×100)

large round or elongated deeply basophilic nuclea and only little becoming the time in the time and the time the cellular bor dess are often indistinct so that it appears as if the nuclea were embedded in a symplysmic mass. The chadow cells stain pale pink with hematoxylin-cosin. They have a distinct border but take noglauclear saining Instead they show a central unstained shadow at the site of the nucleus. In some areas one can see clearly that the shadow cells develop from the basophilic cells (Fig. 190). In tumors of recent origin numerous areas of basophilic cells are usually present. As the lesson ages the number of basophilic cells decreases due to their de velopment into shadow cells and in tumors of long taxading few or no basophilic cells remain. In addition to basophilic and shadow

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cells, areas of cornification are present. The cornification may occur as sheets of horn cells or as small, round centers of cormfication Sheets of horn cells are found only within the masses of shadow cells while the small, round centers of cornification may lie within areas of basophilic cells as well as within the masses of shadow cells. In occasional instances, melanin is present within the tumor The melanın may be located in either the basophilic cells, the shadow cells or the stroma (Turhan and Krainer, Lever and Griesemer)

The stroma of the tumor usually shows a considerable foreign body giant cell reaction adjacent to the masses of shadow cells A frequent but not constant feature is calcification (Fig. 191). The calcium may be present in large sheets replacing the shadow cells or may be pres ent as fine granules within the cytoplasm of shadow cells Occasion ally, ossification of calcified areas takes place (Nicholson, Highman and Ogden) Malignant degeneration does not occur

Histogenesis. The histogenesis of calcifying epithelioma is not established fully Originally described by Malherbe and Chenantais as a calcified epithelioma of sebaceous glands, it has been variously regarded as a tumor of misplaced rests of sebaceous gland epithelium (Sutton and Sutton), as a basal cell epithelioma with degeneration instead of differentiation of cells (Fink, Muehlon) or as a growth intermediary between epidermal cyst and basal-cell epithelioma (Côte) Turhan and Krajner expressed the opinion that calcifying epithelioma arises from that matrix cells. They believe that the baso philic cells are hair matrix cells and that the formation of shadow cells represents a form of keratinization, an attempt at hair shift for mation. In support of their theory, these authors point out that the basophilic cells resemble hair matrix cells, that the small, round cen ters of keratinization observed in the epithelial formations of these numors resemble cross sections of hair shafts and that the melanin in some of these tumors is analogous to the melanin found in normal hair Highman and Ogden have arrived at similar conclusions

Lever and Griesemer regard the basophilic cells in calcifying epi thelioma as primary epithelial germ cells with a tendency to differen tiation into keratotic hair cells. The basophilic cells, thus, are as sumed to have a function akin to that of hair matrix cells, but, being less mature than hair matrix cells, they produce not hair but irregu lar masses of immature hair cells (shadow cells) Accordingly, in a classification of primary epithelial germ tumors, calcifying epitheli oma of Malherbe can be placed under those with differentiation toward hair structures (see Table 6, page 319)

Differential Diagnosis. Apadermal and sebaceous cysts may re

semble calcifying epithelioma of Malherbe if their contents has un

dergone partial calcification and if they have suptured and due to the resulting foreign body giant-cell reaction, have undergone par tial disintegration. They then share with calcifying epithelioma the presence of stregular islands of epithelial cells of areas of calcifica tion and of a foreign body giant cell reaction. However Xbasophilic cells and Ahadon cells are not found It should be remembered that old lesions of calcifying epithelioma may show no more basophilic cells so that then the presence of shadow cells alone is the decisive factor in favor of calcifying epithelioma

Differentiation of calcifying epitheliona from sebaceous cyst with secondary basal cell epithelioma also rests largely on the presence of shadow cells since the areas of basophilic cells in calcifying epi thelioma resemble basal-cell epithelioma, however, in colciffing epithelioma the epithelial structures never show Kalisading of the peripheral-cell layer and often there are areas in which one can ob

serve the transformation of basophilic into shadow cells

BASAL-CELL FPITHELIOMA

Basal-cell epithelioma may occur anywhere on the skin except on the prims and the soles However, the face-particularly the peri orbital region-is by far the most common site of location. The Anucous membranes are never affected. Although basal-cell enthe lioma occurs usually as a single lesion multiple lesions are not infre quent la occa on

present n

types of I c some occur (1) nodulo-ulcerative basal cell epithelioma including rodent ulcer, (2) pigmented basil-cell ept thehoma (3) morphea like or fibrosing basal-cell en et al ana and (4) 11

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" with its size and undered

ing ulcer sur rodent ulcer

ennabileerman 1

Pigmented bisal-cell epithelioma differs from the nodulo ulcera tive type only by the dark pigmentation of the lesion

Morphea like or fibrosing basal-cell epithehoma manifests uself as an only slightly elevated firm yellowish plaque with ill-defined borders over which the skin remains intact for a long time before finally-ulceration occurs

threadlike, pearly border The patches usually show small areas of superficial ulceration and crusting, in addition, their center may show smooth, atrophic scarring

Basal cell epitheliomas cannot be produced perimentally like squamous cell carcinomas. However, they can develop following pro longed administration of morganic assemic (Anderson) (see page 155) and, in rare instances, in areas of radiodermatus (Anderson and Anderson) (see page 122). Furthermore, basal cell epitheliomas of the face occur much more frequently in areas with much sunshine, as in Australia and in the south of the United States, than in less sunny regions, such as England and the porth of the United States.

Basil cell epitheliomas do not need to the Officed Settes
Basil cell epitheliomas do not need to the seems, how
ever, that exceptions to this rule occur (Montgomery, 1928, Foot
Amersbach, Lattes and Kessler) Lattes and Kessler found a toral of
20 cases of metastasizing basal cell epitheliomas reported in the litera
ture Yet many authors doubt the occurrence of metastases in basal
cell epithelioma. Walther expressed the belief that cases reported as
metastasizing basal cell epithelioma represent wrong diagnoses

Histogenesis. Several theories exist regarding the origin of bisal cell epithelionia. Krompecher, the original describer of bisal-cell epithelionia stated in 1903 that he regarded these tumors as area nomas of the bigal cells of the epidermis. He believed that those tumors which showed a tendency to gland formation mutated the embryonal gland formation of the bisal cells. Krompecher's view is still adhered to by many (Montgomery 1940, Teloh and Wheelock). However, Krainz believed that only those bisal cells which develop into glandular cells gave rise to bisal cell carcinomia. Geschickter and Koehler shared this view. They suggested the designation appendage cell carcinomia Mallory, and Haythorn held the opinion that bisal cell epitheliomis were carcinomas of hair matrix cells.

Poot expressed the view that bisid cell epitherioin is were care nomas which develop from distorted primordry of dermal industry rather than from ordinary epidernial basid cells. He street that the tumors took origin from any or all three types of adnessil primordia i.e., hair, sebuceous gland and swert gland and insurated their em

bryonal development

The first author to express doubts that bisid cell epithehomis were carenomis was Adamson who in 1911 stated that, in his opinion bisid cell epithehomas were nevoid tumors originating from latent embryonic fort agoised from their dormant state at a later period of life. He believed that the latent embryonic fort usually were embryonic pilosebaceous follicles but occasionally were embryonic sweat ducts. Several authors have since reached sit.

cently Vallace and Halpert have suggested that basal-cell epithe liomas are benign tumors either of the hair matrix or of the hair palage and have proposed the term trichona for them

unlage and haye proposed the term trichoma for them
It is the puthor's belief that brash-cell epitheliomas are not car
cmomis and are not derived from bisal cells but instead are nevoid

entiated anaphsise cells. In accordance with the potentiality inher ent in the primary epithelial germ to differentiate into sebaceous glinds—sporting glinds and hair differentiation in brisal cell epi thelhomas can be either toward sebaceous gland apocrine gland or hair structures (See Instogeness and classification of epidermal tumos/ pag-317 and Table 6 page-319).

mkus ecenti, irus suggested that basal cell epitheliomas arise not from antémbryonal rest such as the primary epithelial germ but from plurpotenusi cells which base formed during adult, life and crin like the primary epithelial germ differentate in the direction of sebaceous glands apocrine glands and hair. The fact that basal cell epithelioma occasionally arises in areas of radiodermatus and following the ingestion of arisetic is cited by Pinkus as evidence (although to could as well be assumed that these agents merely stumi lite, dormant primary, epithelial germs to active growth).

flueck has pointed out that in basal cell epitheliomas the connective tissue strong always shows a close relationship to the tumor and proliferites with it just as it does in benign fibro epithelial tumors and in adenomas. On the other band in carcinomas the connective tissue usually is stretched then tears until it finally disintegrates. He therefore regards basal-cell epithelioma not as a care

noma but as a solid adenoma

Histopatholog. The characteristic cell of basal-cell epithehoma has a large oval or clonguted deeply basophilic nucleus and little feetbass. The cytoplasm is often defined poorly so that it may appear as if the nucleu were embedded in a <u>symplasmic mass</u>. The nucleu resemble those of the basal cells of the epidermis very closely but the cells of large at a large state of the large state. Its by not has

true car

tint

t₁ comma is an invasive tumor in most instances

Scattered islands of tumor tells are often found away from the main

tumor deep in the decrease and even in the sale.

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latter showing differentiation toward primary epithelial germ structures—i.e. toward sebaceous glands apoctine glands of hair. A sharp dividing line between the two groups cannot be drawn because many undifferentiated basal cell epitheliomas show differentiation in some areas and most differentiated basal cell epitheliomas show areas lacking differentiation. Correlating the finical with the histologic classification it can be strued that the fodulo ulcerative type of basal-cell epithelioma may be differentiated or undifferentiated whereasy figmented basal-cell epithelioma, fibrosing basal cell epithelioma and superficial basal cell epithelioma usually are undifferentiated. If the nodulo ulcerative basal cell epithelioma is it shows no differentiation it is called solid basal cell epithelioma if it shows differentiation it may be either a cystic—in addition of a keratotic basal cell epithelioma.

I UNDIFFERENTIATED BASAL CELL EPITHELIOMAS The group of undifferentiated basal cell epitheliomas includes the solid basal cell epithelioma the pigmented basal cell epithelioma the morpher like or fibrosing basal cell epithelioma and the superficial basal cell epithelioma

A SOLID BASAL CELL PRITHFLIONA (primordial type of basal cell cpithelioma Toot) In this form variously sized and shaped masses of tumor cells are embedded in the dermis (Fig. 192). Some of the masses originate from the surface epider a long connections with it even on serial sections sees a mass of tumor cells originate from licle. In the masses of tumor cells the most of cells often show palisade arrangement, whereas the muclei of the

cells inside lie in haphazard fashigh

A mild or moderately severe inflammatory reaction may be pres ent in the dermis particularly in the more rapidly growing tumors but it may be entirely lacking The connective tissue frequently pro liferates with the tumor and is arranged in parallel bundles around the tumor masses so that a definite mutual relationship seems to exist between the parenchyma of the tumor and its stroma (Hueck Pinlus) The connective tissue nevrest to the tumor masses often indergoes mucinous de enertion Since mucin shrinks during fixa-tion the stroma frequently retricts from the tumor islands so that in prepared sections the tumor islands seem to lie free in cavities (Fig 201) Although this retriction represents merely in artefact caused by shrinkinge during fixation it is quite typical of basal cell epithelioma and aids in il cell epithe squamous cell carcinoma ulcer) lioma often is atrophic.

vome basal-cell eputheliomas with intile or no structural differential tion toward the cutaneous appendages show evidence neverthes of cellular differentiation by presenting two types of cells of cell thas small, elongitud, deeply basophilic nuclei, the other has



primordial basalcell en bas

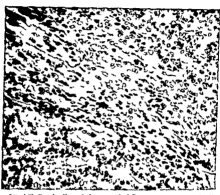
large to mil --

o - and secretory cells of either sebaceous or

apocrine glands

a PIGNENTED BASAL-CELL EPITHELIONA Although the presence of melanin can be demonstrated by ulter-tiains in many basal-cell epitheliomas (Becker) large amounts are encountered only rarely. The

presence of melanin in basal cell epitheliomas can be explained by the fact that melanin producing melanocytes are present not only in the surface epidermis but also in the primary epithelial germ (As the primary epithelial germ matures into the hair these melanocytes arrange themselves among the han matrix cells overlying the hair papilla and form the melanin of the hair (See Lig 9)



IK 133 Basal cell epithelioma vith differentiation into tvo types of cells. One type of cell I as large, ovil, pale in clea, the other type has small elongated dark nuclei (×400)

Basal cell epithchomas with large amounts of piament contain interspersed between the tumor cells numerous melanin laden melanocytes (Bloch). The connective tissue of these tumors contains numerous melapophores (Eller and Anderson Becker Toot)

C MORPHEA LINE OR FIBROSING BASALCELL FLITHFLIOMA In this variant Connective tissue proliferation is much greater than in the other types of bisal cell epithelioma Imbedded in a dense fibrous stroma one observes innumerable groups of closely picked tumor cells arranged in elongated strands (Fig. 194) (Caro and Honell) Most of the strands are small but others are larger and show branch ing The groups of tumor cells often extend deeply into the dermis

D SUPERFICIAL BASAL CELL EPITHELIONA This type of basal cell epithelioma originates from the epidermis in multiple foci (Fig. 195)

The peripheral-cell layer of the budlike proliferations usually shows palisade arrangement. There is littlef penetration into the dermis It may be pointed out that the epidermal buds often show great resemblance to the primary epithelial germ buds as seen in the em



Fig. 191 Mary healthe or fibrosing basal cell epithelioma Innumerable small groups of closely packed tumor cells many of them arranged in elongated strands are embedded in a dense fibrous stroma (x100)

hnorni skin (see Figure I). The overlying gendermis usually shows activities and papillomators. A mild to moderate vinount of inflammatory inflittate composed of lympho ever indignisms cells is present in the upper dermis (Wise Mont gomery 1929).

Some superficial basal-cell epitheliomas after having persisted as such for various lengths of time become true-invasive basal-cell epi

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presence of melanin in bisal cell epitheliomas can be explained by the fact that melanin producing melanocytes are present not only in the surface epidermis but also in the primary epithelial germ (As the primary epithelial germ matures into the lair these melanocytes arrange themselves among the hair mature cells overlying the hair papilla and form the melanin of the hair (See Fig. 9)

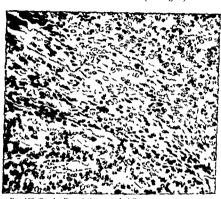


Fig. 193. Based cell epithelioner with differentiation into two types of cells. One type of cell has large oval pule nuclei, the other type has small elongated dark nuclei ($\chi 400$)

Bisil cell epithelionis with Tirge amounts of pigment contain interspersed between the numor cells numerous_melinin_laden melinocytes (Bloch). The Yonnective tissue of these tumors contains numerous inelinopphores (Eller and Anderson Becker Foot).

C MORPHEA LIAT OR PIBROSING BASAL CELL PITTIFITONIA In this virtual (connective tissue proliferation is much greater than in the other types of bisal cell epitheliotin Embedded in a dense fibrous stroma one observes innumerable groups of closely packed tumor cells arranged in elongated strands (Fig. 191) (Caro and Howelf) Most of the strands are small but others are larger and show branching. The groups of tumor cells often extend deeply into the dermis

p superficial basal Cell Epitheliona. This type of bisideell epitheliona originates from the epidermis in multiple foci (Eig. 195)

The peripheral cell layer of the buddike proliferations usually shows paliside arrangement. There is little/penetration into the dermis It may be pointed out that the epidermal buds often show great tesemblance to the primary epithelial germ buds as seen in the em



Fig. 101 Morphea like or fibrosing basal cell epitheli ona Innumerable small groups of closely packed tumor cells many of them arranged in elongated strands are endedded in a dense fibrous stroma (X100)

beyond skin (see Figure I). The overlying Epideemis usually shows acceptly but may slow accutious, and negationators. I mid to moderate amount of inflammatory inflience composed of lymphocities and plasmicells in pressure the upper dermis (Wise, Mont gomer, 1929).

Some superficial basalcell epitheliomas after having persisted as such for various lengths of time, become true-invasive basal cell epi

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theliomas Since this change may at first be limited to a few areas, representative sections throughout the entire tissue block should be examined

A rare type of superficial basal cell epithelioma is the intra epider mal type. In this type, the epidermis shows acanthosis Embedded in the broadened epidermis one finds multiple, sharply demarcated foci of bright of bright bright of bright countries are the properties of bright of the surrounding epidermis by their strongly brighthic strining (Mont gomery, 1929, Suns and Parker)

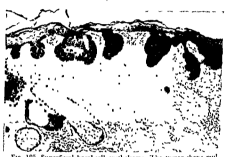


Fig. 195 Superficial basal cell epithelioma. The tumor shows multiple points of origin from the epidermis. Note the similarity between the tumor buds in this illustration and the primary epithelial germbuds in the embryonal skin shown in Figure 1 (X100).

2 DIFFERENTIATED BASAL CELL ENTITIFLIONIAS Differentiation in basal cell epitheliomas may proceed in three directions toward sebaceous glands apocrine glands or hair (Foot, Lever) Those with differentiation toward vebaceous glands are called cistic those with differentiation toward vebaceous glands are called denoted, and those with differentiation toward value are called elegand, and those with differentiation toward value are called elegands, and those with differentiation toward value are called elegands, and those toward in most differentiated basal cell epitheliomas, differentiation is directed toward more than one of the three structures for example, areas of keratinization may be found in a tumor which also shows glandular structures

A CISTIC BASAL CELL EPITHELIONA In this type, cystic spaces are present in the center of the tumor masses. The cysts my form in three ways by degeneration of strome that has become enclosed into the tumor, by degeneration of the center of the tumor masses, and

by differentiation of the cells in the center of the tumor masses to

the victivity of the cyst are apt to be vacuolited (Foot) or even foamy (Pierard and Dupont) (Fig. 195)



FG 198 Cyst c basal-cell et tihel oma. The cyst has formed by distinte grat on of cells with sebaceous differentiation (×50)

a. Abknow BASAL CLIL EPITHELIONA. This type of tumor shows furnitions suggesting ubbular glandlike structures. The cells are arranged in intertuning strands and radially around ulands of connective tissue resulting in a https://linear.com/documents/bit/ by classics one may find human surrounded by cells which lave to proceed the cells with the cells with the cells which lave to proceed the cells with the cells with

C REPRATOTIC BASAL-CELL ENTITIELIDMA (pilar type Foot) In this type cells with clongard nucles and singhily cosmophitic cytoplasm are seen traceting the times short bands or are arranged nn concentric layers are und paraleratonic or keratotic centers. As pointed out by knowpecher who kere described this type of basal-cell epi

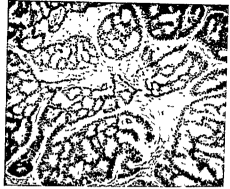


Fig. 197 Adenoid basalcell epithelioma. The strands of epithelial cells show a lacelike pattern. The stroma shows mucoid degeneration (×200)

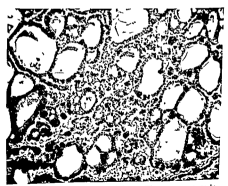


Fig. 198 Adenoid basal-cell epithelioma. The tumor contains lumina surrounded by cells that have the appearance of glandular cells (x200)

Basal Cell Epithel oma

thehoma the elongated cells comify abruptly so that the parakera totic or keratotic centers are sharply demarcated against the sur rounding cells (Fig. 199)

totic or parakeratotic centers in keratotic basal-cell epithelioma de totic or paraseratoric centers in seratoric manacer epitterional de velop from hair matrix cells and represent attempts at hair shaft formation (Foot). The keratinization in keratoric basal cell epithe homa thus some of hair matrix cells which are cells without prickles.

and not one of squamous cells

Some keratotic basal-cell epitheliomas possess large centers of kera tunirition so-called hom costs. (Fig. 200). They are like those observed in tricho-epithelioma (page 363). These hom cysts must not be confused with the horn pearls that occur in squamous cell carci noma (see below under differential diagnosis)

BASAL SOLAMOUS CELL EPITHELIONA Several authors (Darier and Ferrand Montgomers 1908 Juon) have described basal-cell epitheliomas with features of squamous-cell carcinoma. According to Monigomery they represent metamorphosis of basal-cell to squa mans-cell epitheliomis. He stated that from 15 to 20 per cent of all

briajcell epitheliomas present such changes

Tho types of basal squamous cell epithelioms are recognized by the authors named above a mixed and an intermediary type. The nucled type is described as showing partial horn pearly formation with a purplermone ruler than a horns center. The intermediars type is described as showing two kinds of cells those with small clongated deeply basophilic nuclei regarded as basal cells and those with large round pale staining nuclei regarded as intermediaty in character between basal and squamous cells

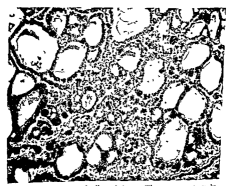
The existence of basal squamous-cell epithelioma is not generally accepted (Leyer Welton Elliott and Kimmelstiel Lennox and Wells It safters that the funced type of basis squamous. Lemma who is used to the the function of the state o

ut secret its cells of either sebaceous or apocrine glands (Fig. 193) see page 373)

MINED CARCINOMA These tumors of one a co-



Fig. 197 Adenoid basal cell epithelionia. The strands of epithelial cells show a Treelike pattern. The stroma shows mucoid degeneration (×200).



F_{1G} 198 Adenoid basal cell epithelioma. The tumor contains lumina surrounded by cells that have the appearance of glandular cells (×200)

However, they ng much more it to remember

that keratinization is not a prerogative of squamous cell carcinoma



tains unusually large horn cysts. (X50)

but occurs also in break-cell epitheliomas with differentiation toward him structures (See Keratoric Basil Cell Ppithelioma," page 373. Keratinization in basil-cell epitheliomas may be partial ("parakera outo centers) or complete ("horn cyste"). The keratinization differs from this seen in the born pearls of symmonis-cell carentomas by occurring alguints without the interposition of gradually keratinizing squirmous cells The laterly common presence in basil-cell epithelioma of areas of retraction of the tumor cell myses from the sur-

Epidermal Tumors

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lesions (for instance, lupus vulgaris, gumma and burns), basal cell epithelioma may stimulate the development of a squamous cell carci noma. Before making a diagnosis of "mixed carcinoma," one must wile out the possibility of pseudo epitheliomatous-hyperplasia occurring in a basal-cell epithelioma (see page 334).

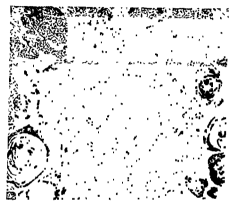


Fig. 199 Keratotic basal cell epithelioma (pilar type of basal cell epithelioma) The cell masses in this tumor are composed largely of clongited cells lying in concentric arringement. Several of the cell masses continua horn cyst in their center. The elongated cells contain novercelles they represent huir matrix cells. Note the altringt and complete keratinization in the horn cysts (X100).

Differential Diagnosis. Differentiation of bisal cell epithelioms from squamous cell carcinoma may be difficult at times—so difficult that many authors have decided that intermediary forms ("bisal squamous cell epithelioma") occur As a rule, however, differentiation is fairly easy. One of the best points of differentiation is that the cells of squamous cell epithelioma stain deeply basophilic, whereas most cells of squamous cell carcinoma at least in Grades I and II, have an eosinophilic tint due to patual, or complete keratinization. In squamous cell carcinoma, Gradex III and IV, the cells, because of

nout type (Hookey) The former type is more common than the lat

12 shows considerable. The acanthosis is due

entitely to upward growth of epithelial cells. Thus, the lower base of the tumor lies above a straight fine that may be drawn from the norm if epidermis at one end of the tumor to the normal epidermis at the other entit and the lesion appears as though tacked on the sur



Fig. 202 Basel-cell papilloms, keratotic type. Low magnification. The superficial location of the tumor is apparent. The lesion appears as it ticked on the epidermis. (X25)

truts of epithelist fells surrounding plands of connective tissue. The horm bare magnates at plues deep into the lesson Because of the normann of these magnations, eyete inclusions of horay material right (1)g 203). They are surrounded by a shell of squamous cells small amounts.

t is located in It is located in Clear cells") are

prominess in the pigmented portions

The follows type of basalcell papilloma shows, originating from the epiderous numerous thin trues composed of a double row of cells. These cells resemble those of basalcell epitheliomas. The tracts rounding connective tissue (see page 372) also aids in differentiating it from squamous cell carcinoma, since such areas of retraction are not found in the latter.

The differential diagnosis between basal cell epithelioma and tri cho epithelioma already has been discussed (see page 364)



Fig. 201 "Mixed carcinoma | basil cell epithehoma (left) and a squamous cell circinoma (right) lie side by side (x50)

BASAL CELI PAPILLOMA (VERRUCA SENILIS, KERATOSIS SEBORRHEICA)

Basal cell papillomas develop, often in large numbers, on the truth, the face and the arms in persons past middle life. They are sharply circumscribed, slightly raised, vertucous, more or less pig mented lesions which often look as if "stuck on" the surface of the skin. The vertucous covering of the lesions tends to have a soft, greasy-consistency. Although most lesions measure only a few millimeters in diameter, an occasional lesion may reach a size of several centimeters.

Histopathology. This tumor represents a squamous cell papilloma in which there are areas of proliferation of cells of the same type as seen in basal-cell papilloma

Basal cell papillomas occur in two types a keratotic and an ade

noid type (Hookey) The former type is more common than the lat

ra shows considerable The acanthosis is due

entirely to upward growth of epithelial cells. Thus, the lower base of the tumor lies above a straight line that may be drawn from the normal epidermis at one end of the tumor to the normal epidermia at the other end, and the leason appears as though tacked on the sur



Fig. 202 Basalcell papilloma keratoric type Low magnification. The superficual location of the tumor is apparent. The lesion appears as if tacked on the epidermis. (X2:)

face of the skin (Fig. 202). The acanthosis is due in part to prolifera

tracts of epithelial cells surrounding islands of connective tissue. The horny layer invaginates in places deep into the lesson. Because of the tortiosits of these invaginations cristic inclusions of horny inaterial result (Fig. 203). They are surrounded by a shell of squamous cells Melan in the mall amounts but to

out c s located in the a

382

rounding connective tissue (see page 372) also aids in differentiating it from squamous-cell carcinoma, since such areas of retraction are not found in the latter.

The differential diagnosis between basal-cell epithelioma and tri cho epithelioma already has been discussed (see page 364)



Fig 201 "Mixed carcinoma" A basal cell epithelioma (left) and a squamous cell carcinoma (right) lie side by side (×50)

BASAL-CELL PAPILLOMA (VERRUCA SENILIS, KERATOSIS SEBORRHEICA)

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Histopathology. This tumor represents a squamous cell papilloma in which there are areas of proliferation of cells of the same type as seen in basal-cell papilloma

Basal-cell papillomas occur in two types, a Leratoric and an ade-

branch and interweave (Fig 204) Considerable amounts of melanin usually are present in the tricis. Cystic inclusions of form, material are absent. The demarcation at the lower border is sharp, just as in the keratotic type of basal-cell papilloma

Although bisil-cell papilloma shows no tendency to invasion of the dermis it may be regarded as related to basal-cell epithelioma because a large proportion of the cells are of the same type as found in basal-cell epithelioma. However actual change of basal-cell papil loma into a frank basal cell epithelioma with deep invasion and ulceration is rare (Filer and Ryan Pinkus)

(For differential diagnosis from nevus verrucosus see page 322 and from ker itosis senilis see page 328)

DERMATOSIS PALULOSA NIGRA

This condition is frequent among Vegroes especially females. The lesions are located on the face especially in the milar regions and consist of minute soft rounded usually hyperpigmented papules

Histopathology The histologic changes are the same as in basal

cell papilloma Melanin pigmentation of the basal layer is pro nounced (Michael and Seale)

CARCINOMA OF SFBACEOUS GLANDS

Carcinomis of sebaceous glands have been described occasionally in the inerature Some authors are more inclined to make this diag nosis than others Warren and Warri for instance reported 28 per sonally observed cases while most other authors have reported only single instances (Beach and Severance)

Circinomas of sebaceous glands occur most frequently on the evelids, originating from merbonian glands which are modified nywhere.

picture ion consists

200) Although Man; cells are undifferentiated distinct sebaceous cells are present Mitotic figures are numerous. Lipid material can be demonstrated not only in the sebaceous cells but also as fine globules in many other cells of the tumor Beach and Severance state that the undifferentiated cells of sebaceous carcinoma differ from those of basal-cell ac 1 1



Fig. 203 Basal cell papilloma keratotic type H gh mag miforation. Thick intervoven tracts of basal cells compose the tumor. Interspersed are cystic inclusions of horny material caused by inaginations of the horny. Type: (X100)

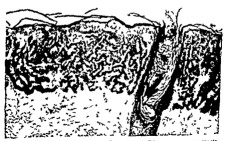


Fig 201 Basal cell papilloma adenoid type. Thin interviewn tracts composed of a double row of basal cells compose the tumor No cystic inclusions of horny material are present (X100)

branch and intervence (Fig. 204). Considerable amounts of melanin usually are present in the tracts. Cystic inclusions of morny maternal are alsent. The demarcation at the lower border is sharp, just as in the keratotic type of braid cell papilloma.

Although basal-cell papilloma shows no tendency to invasion of

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(For differential diagnosis from nexus vertucosus see page 322 and from kerntosis sends see page 328)

DERMATOSIS PAPULOSA NIGRA

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Histopathology The histologic changes are the same as in basal cell papillom Melanin pigmentation of the basal layer is pronounced (Michael and Seale)

CARCINONIA OF SEBACEOUS GLANDS

Carcinomas of vehaceous glands have been described occasionally in the literature. Some authors are more inclined to mike this diagnosis than others. Warren and Warvi for instance reported 28 per sonally observed cases, while most other authors have reported only single instances (Beach and Severance).

Carcinomas of sebaceous glands occut most frequently on the exhibits originating from methomian glands which are modified sebaceous glands (Hagedoom) they may however occur unwhere on the skin (Death and Severance). No charterers of the on the skin (Death and Severance).

20.) Although Many cells are undifferentiated distinct sebaceous cells are present Mitoric figures are numerous Lipid material can be demonstrated not only in the sebaceous cells, but also as fine globules in many other cells of the tumor Beach and Severance strict that the undifferentiated cells of sebaceous carcinoma differ from those of basal cell epithelioma by showing greater variation in size and stape and by having a more acidophilic cytoplasm and a lighter string nucleus.

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It seems that many cases reported as sebaceous carcinoma may be regarded as sebaceous adenoma (see page 347) or as sebaceous epithe lioma, which is a basal cell epithelioma with considerable differentiation of the cells toward sebaceous gland cells (page 353). For in



IN 205 Circinoma of sebaceous glands. Closely picked lobular formations are present in the dermis. They are composed of sebaceous and indifferentiated cells. Some of the latter are applied. (x200).

stance, the five cases of sebaceous carcinoma reported by Savataid as arising in cases of nevus sebaceus are best classified as instances of sebaceous epithelioma

CARCINOMA OF ECCRINE SWEAT GLANDS

Caremomas of the eccrine sweat glands are rare, but their exist ence seems to be established definitely. They do not possess a char acteristic appearance, either clinically or histologically Histopathology. It is inflicult to recognize carcinoma of eccrine sweat glands and to differentiate it from other cutaneous tumors,

carcinomas of sweat glands

forms He pointed out that combinations of these forms were fre quent. He recognized the occasional occurrence of metaplasia of glandular cells into squamous and horn cells.

ADENO ACANTHONIA OF SWEAT GLANDS

Catenomas of sweat glands with glandular as well as epidermal elements have recently been reported under the designation adenoacapthoma of sweat glands (Lever)

Clinically, these tumors resemble squinnous-cell carcinoma. They show shallow central ulceration and may have a vertucous surface. Their site of predilection is the face, especially, the ears



several atypical success glands or ducts are present. (X50)

Histopathology. Adeno acanthomas show tubular and alveolur lumina lined with one or several layers of epithelium (Fig. 206). In areas where the lumina are lined with a single layer of epithelium the epithelial cells have the appearance of glandular cells, but in areas with several layers of epithelium, squamous and partially kera timized cells usually form the inner layers. The lumina are filled with



Fig. 207 Adeno acanthoma of swent glands. High magnification of the atypical sweat glands or ducts seen in Figure 206. Papillary tuffs prototede into their lumin. These swent glands approach in appear time the glandular structures belonging to the tumor (X200).

desquamated cells, many of which are pritially or completely kera timized. In addition, there are solid areas which have the appearance of squamous cell carcinomy. Atypical eccuries weat glands and sweat ducts are present within or at the periphery of these timors (Fig. 207).

Histogenesis. The author originally regarded these tumors as sweat gland carcinomas composed of glandular and epideraral elements and explained the presence of these two structures by the fact that sweat ducts are composed of squamous cells in their epiderimal portion, and of glandular cells in the dermis. It is possible, however, that these tumors are squamous cell carcinomis of alveolar growth in which there is considerable individual cell keratinization resulting in acantholysis in the center of the alveolar formations.

Borelli regards these tumors as dyskeratotic squamous-cell carci nomas which arise from the hair follicle and tend to imitate the structure of glands which develop embryologically together with the follicle

CARCINOMA OF APOCRINE GLANDS

Extramammary Paget's disease represents a carcinoma of apocrine glands (see page 339) No other form of carcinoms of apocrine glands has so far been established

METASTATIC CARCINOMA OF THE SKIN

Cutaneous merastasis of carcinoma is rare with the exception of carcinoma of the breast Next to carcinoma of the breast metastases

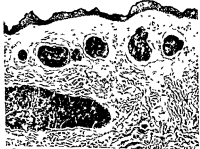


Fig. 208 Metastat c carcinoma of the skin from carcinoma of the breast Inflamn atory care noma. The dermal lymphat es are filed with clusters of t mor cells (×100)

to the skin are observed most frequently in carcinoma of the stomach if e uterus the lings the large intestines and the kidness in that order (Gates). In occasional instances cancer of the new terms of the new t

as seau to cutaneous metastases Dissemination may take place through the lymphatics or through

the blood stream. In carcinoma of the breast cutaneous metastases

Histopithology Adeno aconthomas show tubular and alveolar lumina lined with one or several layers of epithelium (Fig. 206). In areas where the lumina are lined with a single layer of epithelium the epithelral cells have the appearance of glandular cells but in areas with several layers of epithelium squamous and partially kera timized cells usually form the unner layers. The lumina are filled with



Fig. 907. Adeno reanthoms of swest glands. High magnifection of the stypical sweat glands or ducts seen in Figure 406. I spillary tufts protected into their lumin. These si est glands approach in appear since the glandular structures belonging to the tumor (X200).

desquamated cells many of which are partially or completely kera tinized. In addition there are solid areas which have the appearance of squamous cell carcinomy. Atypical eccenic, sweat glands and sweat ducts are present within or at the periphery of these timors (Fig. 207).

Histogenesis. The author originally regarded these tumors as swert gland carcinomas composed of glandular and epidermal elements and explained the presence of these two structures by the fact that swert ducts are composed of squamous cells in their epidermal portion and of glandular cells in the dermis. It is possible however that these tumors are squamous-cell carcinomas of alveolar growth in which there is considerable indusdual cell keratinization resulting in acantholysis in the center of the alveolar formations.

In Afflammators careinoma the skin of the affected breast and the adjoining areas present erythema and diffuse edema simulating ery speks. In Affangectatic extensiona the skin contrins numerous purplish papules and femorrhagic pseudosesicles resembling hemo lymphangioma. In Sancer en curiasse the skin of the breast affected by the extensional and other also the surrounding skin shows dif



Fig. 210 Menantic categories of the skin from carcinoma of the breat Cincer on Currict of thursted area Only few tumor cells are present. Jet e embedded fet see no clings buildes it single row, the state of their amoil number and size if ey may be overlooked as the control of their amoil number and size if ey may be overlooked.

fuse brawny induration interspersed with nodules and punched out ulcerations

Histopathology In inflammatory caremoma histologic examination of the skin reveals extensive mission of the dermal lymphatics—especially of the subepidermal lymphatics—by groups and cords of tumor cells (Fig. 208) (Taylor and Melter). The tumor cells are similar to those of the primary growth stypical in character with large rounded deeply staming nuclei and moderate amount of cytoplasm. Occasional mouser are seen in these cells. There is marked capillary congestion (which is the reason for the inflamma tory appearance chimcally). In addition one observes edema and a slight perivacular lymphocytic infiltrate in the detrins but negligitary and Case).

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usually are caused by way of the lymphatics. In other carcinomas, cutrineous metastases, as a rule, are caused by dissemination through the blood stream, although occasionally dissemination takes place through lymphatics secondary to local lymph node involvement (Gates).



Fig. 209 Metastatic carcinoma of the skin from carcinoma of the breast Cancer en currase nodular lesson. There is considerable fibrosis of the dermis with small scrittered Blands of tumor cells. Some of the islands show a suggestive glandular arringement of the tumor cells. (X200)

CUTAVEOUS METASTASIS FROM CARCINOMA OF THE BREAST

Three types of cutaneous metastases may occur in carcinoma of the breast inflammatory carcinoma telanguectatic carcinoma, and cancer en cutasse. Two or all three of these types may be present in the same patient. If dissemination of meristases through the lymphatics proceeds/rapidly inflammatory carcinoma results in most instances, and telanguectatic carcinoma in rare instances. If dissemination proceeds slowly, cancer en cutasse eventuates (Taylor and Meltzer)

Histopathology Large and small groups of tumor cells are present throughout the dermis. As a vule no tumor cells are found within Xapillaries onlymphitus. A certain amount of fibrosis is usually present. Signs oxinithmention are absent (Gates)

The mension corcinoma readily can be closufied as either an adenocarcinoma [Fig. 211]. a squamous-cell carcinoma or an undiffer

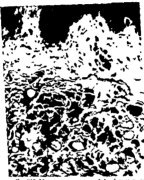


Fig. 212 Metastatic carcinoma of the skin originating from the stomach. Many of the tumor cells are so-called agnet ring cells in which on account of the presence of mucin in the cytoptasm the nucleus is present against the cell wall. (x400)

entiated creanoms. However, it is only occasionally possible to draw conclusions from the histologic appertance of the metastasis as to the site of the primary tumor. For instance, if the tumor cells of the tumoreous internations consimilation the primary carcinoma most likely resides in the gainst metasial rate. The mutin-continuincells present in the metastriss appear, just as in the primary tumor is so-called signet time cells—namely, as large round cells filled with motion which presses the nucleus against the cellular wall. (Fig. 212)

392 Epidermal Tumors

In Clanguectatic carcinoma, the dilated lymphatics contain in addition to groups of tumor cells, varying amounts of erythrocyes (Freeman and Lynch, Leavell and Tillotson). The frequent location of these lymphatics immediately beneath the epidermis causes the clinical resemblance of the lesions to vesicles.

In cancer en cutrasse, the nodular areas usually contain large and small groups as well as sheets of tumor cells lying outside of lym phatics in the dermis and surrounded by fibrosis (Fig. 209). However, the indurated areas often contain only few tumor cells which therefore, may be easily overlooked. The tumor cells are small angular and deeply basophilic and he as single cells in single row lines or small groups between thickened collagen bundles. The arrangement in single row lines, like Indians in a file, is of par ticular diagnostic importance (Fig. 210).

CUTANTOUS METASTASIS I ROM CARCINOMAS OTHER THAN BREAST CARCINOMA

The cutaneous metastases caused by carcinomas of organs other than the breast occur clinically as circumscribed nodules or tuniors. The nodules are usually discrete and moderately firm. Their number may vary from one to several hundred.

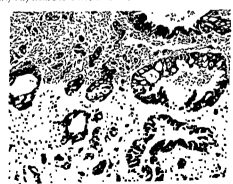


Fig. 211 Metastatic adenocarcinoma of the skin Numeror lumina are present (×100)

Histopathology. Large and small groups of tumor cells are present throughout the detrus. As a rule, no tumor cells are found within Xapillaries or management of the transmission of the cells are found within ent Signs obtiliammation are absent (Cates)

The metastatic carcinoma readily can be classified as either an idenocarcinoma (Fig 211), a squamous cell carcinoma or an undiffer-

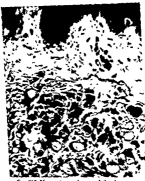


Fig. 212. Metastatic cartinoma of the skin originating from the stomach. Many of the tumor tells are so-called signet ring cells in which on account of the presence of mutin in the cytoplasm, the buckets is pressed against the cell wall. (X400).

entrated caremoma. However, it is only occasionally possible to draw conclusions from the histologic appearance of the metastasis as to the site of the primary tumor. For instance, if the tumor cells of the cutaneous metastasis contain/anocia, the primary caremoma most likely resides in the greator intestinal tract. The mucin containing cells present in the metastasis appear, just as in the neuron as so-called nome.

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19

Mesodermal Tumors

The mesodermal tumors occurring in the skin may be disided into (1) tumors of fibrous tissue (2) tumors of mutood tissue (3) tumors of fatty tissue (4) tumors of nerve tissue, (5) tumors of vascular tissue (6) tumors of muscular tissue and (7) tumors of osseous tissue (6) tumors of nerve tissue under mesodermal tumors is of course not entirely correct but appears justified because mesodermal nerve sheath cells seem to represent an important component of most of the cutaneous tumors of nerve tissue

Benign and malignant tumors occur. The malignant tumors of mesodermal ussue are called sarcomas. They are almost invariably single tumors. Involvement of other organs takes place by way of

metastasis

Gld classifications formerly responsed spindle cell sarconn and round cell sarconn. The terra-spindle-cell sarcoma is spinonjmous with fibrosarconn account cell sarconnas are no longer classified as such but as lymphomas (i.e. stem-cell reticultum-cell lymphodhistic and lymphocytic lymphoma see page 471). Lymphomas differ from sarconnas by their potentially systemic nature which is demonstrated by the fact that frequently they arise in multiple foci. Also the term melanosarcoma is no longer in use unce the mother cell of the tumors formerly so designated is not a mesodermal cell. The preferred term is malignant melanosar.

1 TUMORS OF FIBROUS TISSUE.

DERMATOFIBROMA HISTIOCYTOMA (SCLEROSING HFMANGIOMA NODULAR SUBEPIDERMAL FIBROSIS)

This lesion occurs in the kin as firm indoloni single or multiple nodules. The nodules usually arise in adults and are situated most commonly of the extremites but occasionally elsewhere. Although they are as a rule not larger than a few millimeters in diameter they may measure several centimeters in size. Most lesions have a reddish rolor others are yellowish brown (because of the presence of large amounts of lipid) or blush black (because of the presence

402 Epidermal Tumors

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the rete ridges In some instances, this prohibitation results in bud dung formations resembling superficial basticell epithelioma. The subcutaneous fat is not invaded (Rentiers and Montgomery) Exdence of phagocytic activity is lacking and staining fag from and lipid gues negative results



110. List Dermatchbroma Low magnification The collagers shows disorderly arrangment. Such of the collagen is coung and instead of staming bright red with hematorylin coin stams pade blue and instead of being assem beld in firm bundles, her in individual fibers A moderate number of binoblasts and of appliances are present (250)

Histocytona In thy lesson, in addition to fibriobasts, there are varying numbers of distinctions. They tend to be in nexts Occasion ally the great majority of cells are histocytes. The histocytes are larger cells than the fibriobasts and possess ample amounts of pale typolasm and oxoid rather than so not.

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of large amounts of hemosiderin). In the latter case, the clinical appearance resembles that of a malignant melanoma

Histopathology. Histologically, the nodules can be divided into two types those composed entirely of fibroblasts (dermitofibrom) and those containing, in addition to fibroblasts, varying amounts of histocytes (fustiocytomi)

Various theories exist concerning the histogenesis of this lesion Originally described as dermatofibroma, Moringer, in 1932, proved the presence of phagocytic histocytes in many derinatofibrom's and proposed the term histocytoma for those containing histocytes. He regarded histiocytomas as young dermatofibiomas because of his be hel that the histiocyte represents the parent cell of the libroblast and may develop into a fibroblast (see page 33). Woringer's observations were supported by the studies of Senear and Caro who showed by vital staining with colloidal iron that some lesions which had histo logically the appearance of dermatofibromas contained cells with phagocytic properties and thus were histocytomas rather than der matofibromas Gross and Wolbach, on the other hand, have empha sized the presence of blood vessels in these lesions and have expressed their belief that they represent sclerosing hemangiomas. They regard the cells present in these lesions as proliferating endothelial cells which attempt to form new blood vessels but do not always succeed and instead become engulfed by regressive fibrosis. It would seem however that the development of collagen in these tumors does not represent a regressive process but rather a progressive differentiation Avail different view has been taken by Michelson and by Rentiers and Montgomery These authors believe that histocytomis are not tumors at all but represent a chronic inflammatory proliferation of fibroblasts that may follow trauma. They refer therefore to dermato fibroma and histocytoma as nodular subepidermal fibrosis. The pres ence of histocytes is explained by them as a response to hemorrhige and local tissue destruction

Dermatofibroma. The nodule contains vaying numbers of cells all of which are fibroblasts with spindle shiped nuclei. Much of the collagen produced by these fibroblists is young and instead of staining bright red with hematoxylin cosin it stains a pale blue and instead of being assembled in firm bundles it lies in individual fibers (see Plate 3). The fibroblists and the collagen are arranged in parallel in hands. (Fig. 213). The lesion is

encapsulated, and nearly always

is reparated from the overlying conternits by a narrow band of nor mal collager. The epidermis may be normal or atrophic but com monly shows acanthosis with irregular downward proliferation of

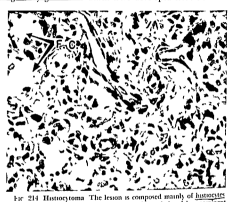
the rete ridges. In some instances this prohiferation results in budding formations resembling superficial bisal cell epithelioma. The subcutaneous fat is not invaded (Rentiers and Montgomery). Evidence of phagocytic activity is lacking and staining for iron and lipid gives negative results.



lagen shows disorderly arrangement Much of the collegen is young and instead of staining bright red with hembled in f

larger cells than the fibroblasts and possess ample amounts of pale choplasm and oxord, rather than spradle shaped nucles (Fig 214). Numerous newly formed Appliances with prominent endothelist cells may be present. The histocytes contain varying amounts of hipid or

hemosiderin or both These substances can often be recognized with out special stuning but are much better visualized when special stunis for iron and lipid are employed The lipid usually is doubly refractile. Some tumois contain a fairly large number of true foam cells and even Touton giant cells. Occasionally, one may find large forcing body, mant, cells which have developed from either histo



There are numerous large and small blood <u>ressels</u> lined by prominent endothelial cells. Several foam cells are present, two particularly large foam cells (F.C.) can be seen in the upper left corner (X400)

cytes or endothelial cells (see below under Nevo xantho endotheli

om1, page 407)

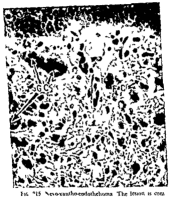
Differential Diagnosis In rare instances, Vermatofibromas show a considerable number of huclei. In such cases, differentiation from fibrosarcoma especially dermatofibrosarcoma protuberans (see page 410) may cause difficulties. However, the circumscribed nature of the lesson especially the absence of invasion into the subcurancous fut the lack of atypicality in the appearance of the nuclei and the absence of mitotic figures, rule out sarcoma (Michelson).

Mistiocytoma with many fat cells closely resembles anthoma tuberosum in a regressive fibrosing stage and may even be indistinguishable from it (Arnold and Tilden). In such cases clinical data, such as the number and the location of the lesions and the

presence or the absence of hypercholesterentia are necessary to arrive at the correct diagnosis

NEVO YANTHO-ENDOTHELIONA

Nevo xantho endothelioma is characterized clinically by the pres ence of a group or groups of sellowish brown nodules arising in early



etc. "12 New santho-endotheliuma The Iesuot is composed mainly of h suocytes Several foam cells (FC) and fore gn-body g ant cells (CC) are present This lesion probably represents a form of h stocytoma (x400)

childhood usually on the extremutes. The lesions involute spon taneously within a few years.

Histopathology Nevo xantho-endothelioma probably represents a young histocytoma Ne m histocytoma flusticoytes predominate mann histocytes have a pale vacuolated cytoplasm and on staining for fit, are shown to contain lipid. Typical foam cells and Touton guint cells are present. There are many capillaries showing prolitera tion of their endothelium. In addition, large foreign body guart cells. 408

are present which originate either from histocytes or from endothe lial cells (Fig. 215)

Schear and Caro and Montgomery and Osterberg regard new vanisho endothelioma as a variant of vanishoma. However, since in patients with nevo vanisho endothelioma the clinical and the blood chemical findings do not support a diagnosis of vanishomatosis, and since the histologic picture does not differ essentially from that of histocytoma, it seems best to regard the resion as a histocytoma New variantho endothelioma differs from districctiona only by showing from cells and ligant cells of the foreign body type in larger number.

RETICULO HISTIOCYTOMA

This rare condition occurs as solitary or multiple, fairly large cutaneous nodules in adults. The lesions may remain stationary or may involute.

Histopathology. The histologic picture is characterized by the presence of large, bicarre shaped, multinucleated giant cells with abundant, pale cytoplasm and vesicular nuclei. They are separated from one another by fibrous connective trisic (7ak.) Occasionally, it is possible to demonstrate the presence of fat within the cytoplasm of these cells. Vital staining with colloidal iron indicates that the large cells have phagocytic properties and are reticulo endothehal in nature (Caro and Senear).

It is likely that reticulo histocytoma is an unusual type of histocytoma. Caro and Senear assume that it is not a true neoplasm but actually a granuloma.

KELOID

Keloids represent a post traumatic tissue proliferation. They are red raised and firm and have a smooth, shing surface. Occasionally clawlike projections radiate from the edges of the lesions.

Histopathology, Keloids cannot be differentiated from a dermuo kuroma ana histologic basis. In an early keloid one linds, in admition to thick, intertwining bundles of collagen a moderate number of fibrollasts (Fig. 216). An old keloid may show but few cellular elements.

HBROSARCOMA

Fibrostroma occurs in the skin in two forms as true fibrosarcoma and as derinatofibrosarcoma protuberans. However, there is justifiable doubt that the latter represents a real sarcoma.

TRUE FIBROSARCOMA

True fibrosarcoma starts only tately in the dermis, more commonly, it starts in the subcutaneous fat (Broders, Hargrave and

Meyerding) The tumor usually feels firm and irregular on palpation and at first is covered by normal stan. It murilly grows quite rapidly and as it increases in size the overlying stan at first shows purplish dissolvation and furilly ulcer iron. Strellite lesions frequently develop. Metastases occur sooner or fiver. They usually spread by may of the blood stream especially in the lungs. The region if lymph nodes are modived only rarely (Genele).

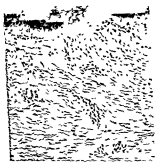


Fig. 216 kelood. The lesson is composed of thick in cruming bundles of collagen. The collagar is estacely maine and strins cosmophilic throughout. I moderate number of nuclei are present therefore this kelood is of relatively retent genesis. (X900)

Histopathology. This tumor is very feelfular. The nuclei vary greatly in use shape and straining qualities. Although most nuclei it is espandis shaped others appear round or otal. Some nuclei it is necessary dark, while others appear residular with irregular chromatur structure. Typical as well as at pixal intoxic figures are present, often in large numbers. The nuclei tend to be in buildes, that extend in various directions bind to not form whorls. It some areas the nuclei fie in dense clusters. Thus, histogracomas present an extremely, disorderly arrangement of the nuclei. Usually, these tumors contain some collagenous tissue in which the nuclei appear to be embedded.

but highly malignant fibrosaryomas may possess little or no collagen Instead, on staining with a Kirculium stain, they are seen to contain numerous reticulium fibers (Foot)

Because of the invasive nature of these tumors, ulceration at the surface and infiltration into the subcutaneous tissue and underlying

structures frequently are observed

Differential Diagnosis. A highly malignant fibrosarcoma with no production of collagen may be difficult to differentiate from squa mous cell carcinoma, Giade IV, amelanotic malignintamelanoma stem cell imphoma or reticultum cell imphoma on thorough in spection, however, one will usually find, invigurmous-cell carcinoma Grade IV, some tendency to keratinization and connections of the tumor with the epiderms (page 331), in milignam melanoma, one will find "junction activity" at the epidermo dermal border (page 438), while insten cell and reticulum cell lymphoma spindle shaped nuclear forms are absent, since all cells are 'round cells ' (see piges 171, 475)

Spindle celled malignant tumors may form in areas of radioderma turs Although they have the morphologic appearance of a fibrostriction, it is likely that most, if not all, represent Grade IV spindle celled squamous cell caremonn (see page 333) (Gentele, Blom Ides)

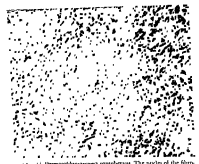
DERMATOFIBROSARCOMA PROTUBERANS

Denmitofibrosincomin protuberans represents a slowly growing tumor which has its origin in the dermits. It begins with one or set

tumor which has its origin in the dermis. Rebegins with one of several closely sets small, hard nodules which are reddish or bluish As the nodules coalesce, they form one or several plaques. On these plaques protruding tumous subsequently develop and may ulcerate Although this tumor is locally invasive as a rule, it does not give rise to whetastases. In exceptional cases, metastases may occur many years after the appearance of the tumor. Binkley, for instance, re yours after the tumor had first appened oost has expressed the belief that dermatofibrosarcoma protuberans was not truly malignant because of its slow rate of growth and the absence of metastases in most cases. He regards the tumor as intermediary between demails fibroma and true fibrosarcoma partaking of the nature of both. He prefers to call it progressive recuirent dermatofibroma.

Histopathology. The histologic appearance of the tumor does not differ materially from that of true fibrosarcoma (Binkley) A differ entiation of the two tumors on a histologic basis therefore is not always possible Just as in true fibrosarcoma the nuclei he in irregu

degeneration of the collagen is observed commonly in parts of the tumor (Binkley). Ulceration at the surface and invasion into the sub-cutaneous fri occur



110. 11/ Dermatohbrosarcoma protuberans. The nuclei of the fibroblasts are arranged in irregular trands and whorls. They show a slight degree of atypicalist. In contrast with miliginatic fibrosarcoma formation of sollagen is well in evidence (X200)

Differential Diagnosis For differentiation from dermatofibroma, see page 406

2 TUMORS OF MUCOID TISSUE

AMOYYM

Uthough fibroms and neurofibroms may undergo more or less complete mucoid degeneration and then present in such degenerated atervilse aspect of myxoma myxomas mry asise as such

Clinically myxomas present themselves as fairly well circum scribed rather soft, usually intracutaneous tumors over which the epidermis is normal

but highly malignant fibrostryomas may possess little or no collagen Instead on staining with a Attendum stain they are seen to contain numerous reticulum fibers (Foot)

Because of the invisive nature of these tumors ulceration at the surface and infiltration into the subcutaneous tissue and underlying

structures frequently are observed

Differential Diagnosis A highly malignant fibrosarcoma with no production of collagen may be difficult to differentiate from squa mous cell carcinoma Grade IV amelanotic malignant intelanoma stim cell hymphom or reticulum cell hymphoma. On thorough in spection however one will usually find involution on cell carcinoma Grade IV some tendency to keraturation and connections of the tumor with the epidenius (prae §31) in malignait melanoma one will find junction actuary at the epideriuo dermal border (page 458) while instem cell and reticulum cell lymphoma spuidle shaped nuclear forms are absent since all cells are round cells (see pages 1711 175).

Spindle celled malignant tumors may form in areas of radioderma titis. Uthough they have the morphologic appearance of a fibrosit coma it is likely that most if not all represent Grade II spindle celled squamous cell caremoma (see page 33.3) (Gentele Blom Ides)

DERMATOLIBROSARCOMA PROTUBERANS

Definition of the production o

Although this tumor is locally invisive as a rule it does not give rise toylietistases. In exceptional cases meristases may occur many seria after the appearance of the tumor. Binkley for instance reported a case in which metastases caused the death of the patient. 38 years after the tumor had first appeared, osta has expressed the belief that derinatofibrosia comp protuberans was not truly malignant because of its slow rate of growth and the absence of metastases in most cases. He regards the tumor as intermediaty between dermation fibronia and true fibrosia comparations of the nature of both. He prefers to call it progressive recurrent demands from

Histopythology The histologic appearance of the timor does not differ materially from that of true fibrosaccoma (Binkley) 1 differ entiation of the two tumors on a histologic basis therefore is not always possible Just as in true fibrosaccoma the fuclei he in irregu

atypicality of the cells favor a diagnosis of myxosarcoma. The cells tend to be stellate and multipolar

SYNOVIAL CYST OF THE SKIN (MYXOMATOUS DEGENERATION CYST OF THE SKIN)

This lesion occurs most commonly on the hands particularly near the terminal interphalangeal joints. The lesion consists of a small semiglobular translucent tumor surrounded by erythema. When punctured a clear mucinous fluid evudes.

Histopathology Histologic examination reveals a cystic cavity in the detrins or the subcutineous tissue. The lining wall is made up of fibrous tissue and shows no endothelial or eputhelial lining. The cyst is filled with mucoid material.

The histogenesis is not clear. Macker and Andrews regarded the lesion as a synovial cyst. Savatard held that the lesion formed because of mucoid degeneration of a libriom? Woodburne sixted that he had never seen any exidence of a previously existing solid tumor. He expressed the belief that the lesion arises because of my comatous degeneration of fibrious tissue of the detrins or of other fibrious tissue.

3 TUMORS OF TATTY TISSUE

LIPOMA

I spomas occur as single or multiple subcutaneous soft rounded lobulated growths which may or may not be movable against the overlying skin. In rive instances multiple lipomas have been reported as present not only in the subcutaneous but also in the vis ceral fat deport. This condution is known as systemic multicentric lipoblasiosis (Tedeschi)

Histopathology Lipomas are composed of fat cells and may or may not be surrounded by a connective tissue capsule. The fat cells in hipomas usually do not differ from normal fat cells. Occasionally they are slightly larger. In some hipomas there is more in others less of a connective tissue framework than in normal subcutaneous fat. In we containing a considerable proportion of connective tissue are called filtrolipomas.

Systemic multicentric lipoblastosis shows not only adult fat cells as tipomas do bui also embryonal fat cells and undifferentiated mesenchymal cells with all intermediate stages. These lesions are hamitiomas inalogous to the lesions of ion Recklinghausens disease and not sarromas. These show no cellular disorder and no mitotic figures (Tedeschi).

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Histopathology. Myxomas contrun embryonal fibroblasts that possess the ability to produce mucin in a minner similar to that of the embryonal fibroblasts of the umbilical cord. The number of these cells in myxomas is usually small. Those present are spindle shaped or stellate cells possessing multipolar processes. The stroma appears homogeneous and gelatinous. It string pale blue with hematoxylin.



Fig 218 Myxoma The stroma appears homogeneous and in some places gelatinous. The empty spaces are caused by shrinkage of the mucin in the process of fivation (×200)

and eosin, and red with Best's mucicarmine stain. Because of shrink age in the process of fixation empty cleftlike spaces frequently are observed in the stroma (Fig. 218).

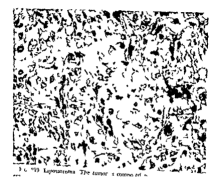
Many myxomas are not pure but contain other mesodermal elements. Thus fibromyxomas and lipomyxomas occur

MYNOSARCOMA

Clinically, mynosurcomps are as a rule primarily subcutaneous tumors which, as they grow in size may cause ulceration of the skin Histopathology. A decision whether a tumor represents a mynoma or a mynosurcomp is often difficult. A high degree of cellularity and

extoplasm (Fig. 219). There is a loose meshwork of connective tissue which usually but not always is mucoid (Stout)

A highly malignant liposarcoma contains highly asspiral lipo blasts t ith bizarre nucles. The amount of fat in them is often small There are areas which resemble fibrosarcoma. The stroma usually is mucoud



MALIGNANT HIBERNOWS

ITe clinical appearance of malignant hibertioma is the same as that of cit er liposarcomas

Histopathology The cells comprising this tumor are rounded, have a centrally placed nucleus and contain numerous small and farge vacuoles of fat in their cytoplasm. Those containing large sacuoles are called mulberry cells. Some of the cells are of very large size and contain multiple bizarre nuclei (Fig. 290). The off

Com

HIBERNOMA

Hibernoma is a rare, benign, solitary, soft, subcutaneous tumor which may slowly increase in size Clinically, it is indistinguishable from lipoma

Histopathology. The tumor is composed largely of multilocular fat cells which resemble those occurring as brown fat in the hiber nating gland of hibernating animals. However, since multilocular fat cells resembling those of brown fat may occur as a transitory phase in the muturation of ordinary, yellow adipose tissue, it is possible that hibernoma merely represents a tumor of embryonal yellow fat cells (Sutherland).

Histologic examination reveals a lobulated tumor composed pre dominantly of closely spaced, round or oxal cells with a distinct cell membrane and centrally placed nucleus. The cells contain either granules or small vacuoles or locules which stain positive with fat stains. This fat is in contrast with mature fat, doubly refrictele to polarized light (Sutherland). In some multilocular cells coalescence of smaller into larger locules can be seen, with eccentric displacement of the nucleus. There are, in addition a few unilocular, normal mature fat cells which are larger than the granular and multilocular cells. There thus seems to be evidence of a transition from granular to multilocular and even to unilocular fit cells (Brines and Johnson).

to multilocular and even to unilocular fit cells (Brines and Johnson)

Differential Diagnosis. The tumor can be differentiated from ma lignant hibernoma by its regular architecture and the absence of mutotic figures and multinucleated cells

LIPOSARCOMA

Liposarcomas as a rule arise as such but they may develop within a pre-existing lipoma (Sternberg). They not only occur in the sub-cutaneous fat but also may be found wherever there is fatty tissue. Those located in the subcutaneous fat present themselves as diffuse nodular infiltrations. Metastases are common, especially to the lungs and the liver.

Histopathology. The histologic picture varies somewhat with the degree of malignancy. Moderately inalignant tumors are easily recognizable as fatty tumors, while very malignant ones may not be identifiable as such unless fat stains are employed.

A moderately malignant liposyncoma consists of adult fut cells and moderately atypical immature fat cells (lipoblasts). The lutter have a spindle shaped or stellate nucleus and contain fat droplets in their

appear cut in transverse and others in longitudinal direction (Fig 221) Each nerve bundle is surrounded by fibrous tissue



Fig. 221 Acuroma Numerous thick bundles of medullated nerves are present. Each nerve bundle is surrounded by fibrous tissue $(\times^{\circ}00)$

NEUROFIBROMATOSIS (NON RECKLINGHAUSEN S DISEASE)

Neurofibromatons is characterized by the presence of multiple quinneous turions which possess a characteristic soft consistency and usually are Besh-colored but may be brownish or isolocous. The lesions are semiglobular or pediunculated and vary considerably in pare Occasionally large pendulous Habby misses tengling a pound on more are encogniered in most cases in addition to the tumors the stan shows fellowish brown pagmented mixules of various size and shape—so-called cafe an last spoots

Histogenesis \text{\text{curofibromas}} are tumors of nerve sheaths \text{\text{Normally}} each neurite or iron whether myelinated or not, is sur rounded by a neuro-ectoderinal sheath the schwannian sheath, and

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Stout regards this as evidence that ordinary adipose tissue and brown fat come from the same ancestral lipoblastic cell

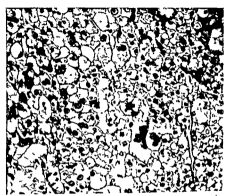


Fig. 220 Malignant hibernoma. The fut cells contain bizarre nuclei and are filled with small and large vacuoles of fut. Those containing large vacuoles are called mulberry cells. There is hardly any stroma (X200).

4 TUMORS OF NERVE TISSUE

Three types of nerve tumors may occur in the skin neuroma neurofibroma and neurolemoma

NEUROMA

Neuromas nearly always occur as single tumors secondary to an injury of a nerve. In rare instances, they occur as single (Duemling) or multiple (Ludy) tumors apparently without preceding nerve in jury Neuromas are small reddish nodules which may or may not be

A painful

Histopathology In their histologic appearance neuromas resemble
the amputation neuromas which represent a hyperplasia of nerves
and are not tumors. For this reason. Ludy preferred to regard also
the nontraumatic neuromas of the skin as neuromation hyperplasia.

Neuromas show numerous thick bundles of nedullated nerves in the dermis extending in different directions. Thus some bundles

It is unfortunate that the schwannian cell and the endoneutral futroblast eatment be differentiated by present histologic methods since the schwannian cell can also produces, etculum fibers and college (Masson, Murry) and Stony) the pressure of these structures in neurofiltromas does not decide the issue, schurzay and Stony believe that their findings in usue cultures favor the schwannian origin of

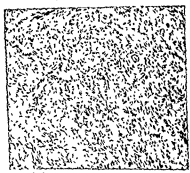


Fig. 223 \eurofibroma (son Recklingbausens duesse) In this tumor mucoul degeneration of the collagen has taken place. This is a not uncommon occurrence in lesions of neurofibroma (\$\infty\$200)

neurohirona because the mode of ourgrowth of schwamman cells from normal nerves and of cells in neurofibroma is very similar

Histopathology Histologic examination of the cutaneous tumors shows them to be well incumerated but not/encapsulated. The tumor mayers often extend into the subcumberous far They are composed observes from the tumor between the total product of the cutaneous strangement of the fibrils is characteristic of neutrofibrious (Fig. 222). With hematoxylin

Mesodermal Tumors

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a mesodermal sheath, the endoneurium (see page 22). It is still a matter of controversy whether the cells composing neurofibromas represent neuro ectodermal schwannian cells or mesodermal endoneurial cells. Whereas you Recklinghausen originally regaded the tumors as derived from the connective tissue of nerves, Verocay, in 1910, first suggested that the cells of the tumors were immature cells.

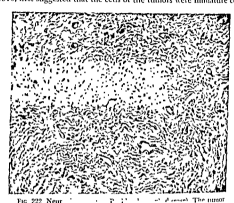


Fig. 222 Neur P descent The tumot is composed of gether and tend

pale blue with hematoxylin eosin (×200)

of the sheath of <u>Schwann and thus neuro crodermal</u> cells. He suggested that the tumors be called neurnomas. Among more recent writers on the subject, some (Murray and Stout) subscribe to Vero cay's view. Others (Penfield, Tarlov) still adhere to the view that the tumors develop from the perineural connective tissue. Many store vecent authors however, feel that probably both ectodermal

rey) Masson, who observed that neural elements were considered in young tumors but sparse or absent in old tumors concluded that neurofibromas basically were neural tumors! However, with aging the neural elements gradually degenerated and were replaced by connective tissue proliferation.

nervous systems and of the bones, however, it is a not uncommon occurrence Hosoi, in 1931, found 65 instances of malignant degeneration reported in the literature; he stated that this represented 13 per cent of all the reported cases of neurofibromas. The conclusion drawn by Charache that 'sarcomatious transformation is present in 13 per cent of all cases of multiple neurofibromas' is not justified, however, because in neurofibromatosis, as in other common diseases,

. . .

fibroma In some, however, the pattern of a neurofibroma is pre served there is still a wavy arrangement of the collagen fibrils, but the nuclea are uncreased in number and are atypical (Fig. 224) Nerve yfibers are never found (Stout, Wachstein and Wolf)

NEUROLEMOMA

Neurolemoma occurs as either solitary or multiple tumors but not as adjustenatured disease like neurofibromatosis. The lesions which are asymptomatic are found in the dermis or in the subectuanceous ussue. They can reach a diameter of several centimeters and undergo

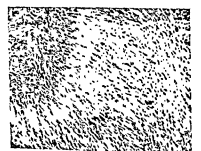


Fig 225 Neurolemona Numerous elongated nuclet are arranged in a streaming (subion In the center of the field lies a so-called verocay body formed by a double pulsade of nuclei enclosing a space nearly decord of nuclei (x200)

uniform in size and rather pile staining Listic fibers are absent in the tumors. On staining with Foot's stain numerous wavy reticulum fibers are seen. In whost neurofibromas a few nonmedulated thin long nerve fibers can be recognized on struing with special nerve stains, such as Bodan's stain (M. Naux and Montgomes).

Not infrequently which degeneration of the collagen is observed in parts of a tumor or in an entire tumor. In such cases, the nuclei

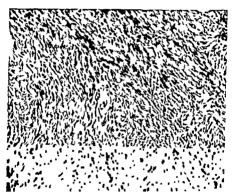


Fig 224 Neurofibrostreoma (von Recklinghausen's disease) The wavy pattern characteristic of neurofibroma is preserved but the nuclei are increased in number and are atypical (×200)

are embedded in a homogeneous <u>pale blue</u> ground substance (Fig. 223). One must be familiar with this change because it results in a histologic picture quite different from that usually associated with neurofibright

The pregnented muculu lessons seen in neurofibromatosis the so called cafe au lut spots merely show increased melunin pigmenta

tion in the basal layer

Systemic Lesions There may be tumors of the central and periph eral nervous systems of the viscera (Grill and Kuzma) and of the bones Involvement of a bone may lead either to extensive destruction or to hypertrophy of bony tissue (Westcott and Ackerman)

tion or to hypertubility of Malignant degeneration of neurofibro mas of the Kim is rate. In tumors of the entral and the peripheral

Nexus Nammeus is characterized by one or several dull red or bluish red patches of irregular outline, nonelevated above the level of the skin //

Nevus vasculosus is a raised, bright red, soft, often lobulated tumor. When lobulated, its appearance resembles that of a strawberry,

Ravernous hemangioma consists of a large, soft, subcutaneous mass If, as is often the case, the lesion is located on the face, consid-



lated apillaries with considerable proliferation of their endothelial cells

erable deformity may result. The overlying skin may be possed to

con a present (Fig. 22b)

NEVUS VASCULOSUS, which represents a capillary hemanorom addition to -- cons

prol

around capitally lumina. In addition, they form solid strands and masses with little evidence of vascular lumina

CAIFRAGES HEMANGIOMA shows in the lower dermis and in the subcutaneous tissue large, irregular spaces filled with blood. They Mesodermal Tumors

partial cystic degeneration Malignant degeneration does not occur, however.

Histopathology. The tumor arises from the sheath of Schwann of a peripheral nerve and is composed of schwannian cells. It is well encapsulated One observes numerous bands of closely spaced, elon gated nuclei arranged in a twisting, streaming fashiom Here and there, the nuclei he in two rows or palisades enclosing a space nearly devoid of nuclei Such formations are called Verocay bodies (Fig 225) In addition to these areas of streaming and palisading there are areas in which the schwannian cells are embedded in a loose mesh work of fine connective tissue fibers. Small cysts may be present which by coalescence may form gross cystic spaces (Stout) No nerve libers are found in neurolemonia. Mast cells often are present in conspicu ous numbers

5. TUMORS OF VASCULAR TISSUE

HEMANGIOMA

Hemangiomas may be divided into three clinical types (1) nevus flammeus (port wine nevus), (2) nevus vasculosus (strawberry muk) and (3) angioma cavernosum

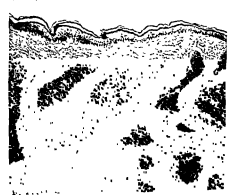


Fig 226 Hemangioma (nevus flammeus) The capillaries are in creased in number They are dilated engorged with blood and lined by only a single layer of endothelial cells (X100)

News Hammeus is characterise for the or even and the blush red patches of arrealize realize, none-critic acree the erec of the skin //

Nevus aboulosus us a rases, briefe res, sons orea horaster rumer.

When lobulated, its appearance sembles that if a start e . Lavernous herrargirum consess et a arms sen surcinnerens mass. If as is often the case, the feather is ordered in the take the pro-



Fig. 227. Hemangoma (nerus vasculosus). There are numer as distinct apillaries with considerable productation of their releases at the considerable productation of their releases at the considerable productation of their releases at the considerable products and their releases at the considerable products are their releases at the considerable products at the considerable products are the considerable products at the c

erable deformity may result. The overly word - -

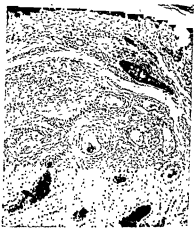
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NEVUS VASCLEONES which represents a car to - - e oo ubha

masses with little evidence of vascular lumina HER they form solid strands and

CAVERNOLS HEMATCHOMA theses in the lower dermis and in the subcutaneous tissue large, priegular spaces filled with blood. They are lined by a single layer of thin endothelial cells and by thick walls (Fig. 228). The thickening is produced mainly by overgrowth of advantial cells

Sclerosing hemangioma, since it is a dermatofibroma tather than an angioma, is discussed under "Dermatofibroma" (see page 403)



Fit 228 Hemangioma (hemangioma cavernosum). The blood vessels show considerable thickening of their walls produced by overgrowth of adventitial cells (×100)

GRANULOMA PYOGENICUM

This lesson, which usually is single, consists of a dull red, soft or fleshy, raised, more or less pediunulated nodulg. Its size varies from 0.5 to 2 cm in diameter. The surface may show a smooth, atrophic epidermis but often is covered by crusts. The lesson bleeds easily when traumatized.

Histopathology. On histologic examination, one finds a circum scribed, raised, peduncul ited lesion covered by a flattened epidermis and containing numerous newly formed capillaries showing varying

degrees of dilatation (Fig. 229). A slight to moderate amount of endo thehal proliferation is usually present. The capillaries are embedded in a loose, edematous occasionally mucoid connective tissue. At the neck of the pedunculated lesion the epidermis usually shows acan thotic inward growth thus forming a so called epidermal collarette



. Giannioma pyogenicum The lesion is pe dunculated it is composed of numerous capillaries em bedded in a loose edematous stroma No inflammatory reaction is present (X50)

In early lesions one finds no dallammatory reaction (Freund) In older lesions because of the fact that the thinned epidermis usually erodes secondary inflaminatory changes are often present in the stroma and may give the tumor a granulomatous appearance

caused b

ture favo

... cand has pointed out that, even when the lesion shows a marked granulomatous reaction, the center of the Differ

which a

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of the raised pedunculated growth of the lesion, the presence of edema or mucoid degeneration in the stroma, the thinning of the epidermia over the tumor and the collarette formation of the epidermia at the pred of the tumor.

ANGIOKI RATOMA MIBLLLI, ANGIOKERATOMA CORPORIS DIFFUSUM

In angiokeratoma fibelli, one observes on the dorsum of the fingers, on the toes and on the knees from pinhead to bean sized dark-red, vascular papules with a vertucous surface. The disease arises in young subjects.

In angiokeratoma corporis diffusum, dark red, vascular papules are present in great number over the entire trunk. They are smaller and show less/hyperkeratosis than those of angiokeratoma Mibelli Angiokeratoma corporis diffusum may be associated with internal lessons such as swelling and vacuolation of the muscle fibers of the blood vessels and of the heart (Ruiter)

SENILL HEMANGIOMA

Semile hemangiomas are small, raised soft, dark red nodules, mers uring usually between 1 and 3 mm in diameter. They may be present in large numbers in persons past middle life. Their sites of predilection are the face and the upper furns.

Histopathology. Semile hemangromas, within a circumscribed area of the upper dermis show numerous dilated capillaries lined by a single layer of endothehum. The spidermis shows futtening of the rete ridges (Beek). Occasionally, however they may show acanthosis as well as hyperkeratosis and even inclusion of dilated capillaries into the epidermis. In that case, histologic distinction from angiokera toma may be impossible (1 raub and Tolmach).

NEVUS ARANEUS

Nexus araneus or spider nexus is characterized by a central slightly elevated red dot from which fine blood vessels radiate. Oc casionally pulsation can be observed Spider nevi are common on the upper half of the face

Histopathology According to Walsh and Becker, nevus araneus represents a small arterior enous anastomosis. They concluded from serial sections that in this lesion an arteriole ascends high into the dermis where it changes directly into a vein. The vein divides into a network of smaller venules. The latter give the lesson its spiderlike

appearance clinically

Ho * ** *** differ in their composi nes of vascular arrange tion al vessel of the spider ments

nevus" to be an artery which branched successively into arterioles and capillaries. In the other type the so-called glomus type they found the afferent artery to be connected with the central vessel of the spider nevus by a short junction which had the histologic up pearance of the Sucquet Hoyer canal as seen in the cutaneous glomus In contrast with the usual glomus however the central vessel of the spider nexus was not a collecting vein but continued into estrelleges

OSLER'S DISFASE (FAMILIAL HEMORRHAGIC TELÀNGIECTASIA)

This familial disease is characterized by the presence of numerous telangiectases on the skin and the mucous membranes. The presence of telanguectuses on the mucous membranes may result in Viemor thages from the nose the mouth, the stamach the kidney the rectum or the yagina

Histopathology Scattered greatly dilated capillaries are present in the upper dermis. Fingerland and Janousek noted that the venules in the lower dermis showed narrowing of their lumina and an in crease in the number of adventitial cells

LA MPHANGIOMA

I superficial and a deep variety of lymphangioma exist. The super ficial variety lymphangioma circumscriptum is characterized by the presence of groups of small thick walled vesicles resembling frog s spawn Some of the vesicles may show a verrucous surface

The deep variety lymphangioma cavernosum, causes diffuse en

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of the raised pedunculated growth of the lesion, the presence of edema or mucoid degeneration in the stroma, the thinning of the epidermis over the tumor and the collarette formation of the epidermis at the neck of the tumor.

ANGIOKERATOMA MIBELLI, ANGIOKERATOMA CORPORIS DIFFUSUM

In angiokeratoma [Mibelli, one observes on the dorsum of the fingers, on the toes and on the knees from pinherd to bean sized dark red, vascular papules with a verrucous surface. The disease arises in young subjects.

In angiokeratoma corporis diffusion, dark red, vascular popules are present in great number over the entire trunk. They are smaller and show less hyperkeratosis than those of angiokeratoma Mibelh Angiokeratoma corporis diffusion may be associated with internal lessons such as swelling and vacuolation of the muscle fibers of the blood vessels and of the heart (Ruiter)

Histopathology. Histologic eximination of the cutineous lesions reveals in both diseases (elangiectasias of superficial location asso crated with changes in the epidermis consisting of hyperkeratosis, acanthosis, irregular proliferation of the red ridges and papilloma tosis. Greatly diluted vapillaries lined by a thin layer of endothelial cells he in the papillae and are partly or completely surrounded by the hypertrophic stratum malpighi. If completely surrounded the dilated capillaries have the appearance of intra-epidermal black cysts. Some of the blood cysts may have lost their endothelial lining Atrophy of the stratum malpighi may occur directly over blood cysts because of the pressure which the cysts evert on the overlying epidermis.

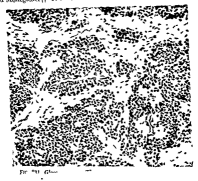
SENILE HFMANGIOMA

Senile hemangiom's are small traised soft, dark red nodules, mers uring usually between I and 3 mm in dameter. They may be present in large numbers in persons past middle life. Their sites of predilection are the face and the upper turns.

Histopathology. Semile hemangiomas within a circumscribed area of the upper dermis show numerous dilated capillaries lined by a single layer of endothelium. The spidermis shows flatening of the reteridges (Beck). Occasionally however they may show acanthosis as well as hyperkeratosis and even inclusion of dilated capillaries into the epidermis. In that case, histologic distinction from angiokera toma may be impossible (Traub and Tolmach).

GLOMUS TUMOR

This tumor usually occurs as a single small, deep seated pink or purphish nodule which is tender and gives rise to severe paroxysmal pains. In rare instances, there are numerous lessons in which case tenderness is absent in most of them (Wedman and Wise, Eyster and Montgomety). The most common sites of the solitary lessons are



a senual venel (X200)

the mad hed and the fingertips however, the lesions may occur else where

Hinopathology The glomus tumor represents a benign tumor of the cytaneous glomus a structure composed of an atterial segment the Sucquet Holyer ranal and a senous segment. The normal Sucquet Holyer canal possesses a nation lumen limed with a single base of endothelist cells and a Mach matele of glomus cells (see page 24) choints cells have a family cosmophilic extendes.

ej fil

A glomus Yumor is composed of Yascular lumina and numerous

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largement of the affected region for instance macrochella and

Histopathology In lymphangional circumscriptum one observes in the uppermost portion of the dermis cystically directlymph vessels lined by a single layer of endothelium (Fig. 230). They contain

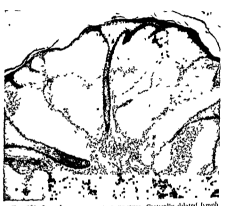


Fig. 230 Lymphangiona circumscriptum. Cysucilly dilated lymph vessels lined by a single layer of endothelial cells are present in the upper dermis. The ep derm s shot s down and growth and more or less surrounds some of the lymph vessels. There is moderate hyper ferators (750)

coagulated lymph and lymphocytes. The stratum malpighu varies greatly in thickness. Over some of the lymph cysts it is thinned else where it may show considerable it authors and irregular downward growth. Some of the dilated lymph vessels may be completely sur rounded by epidermis—flyperket itosis is common. The histologic picture may be similar to that of inglokeratomal except that the dilated areas contain lymph fluid instead of blood.

In lymphangioma Aremosum large lymphfilled cystic spaces lined by a single layer of endothelium are present in the dermis and the subcutaneous tissue. There is concomitant hypertrophy of the Aconnective tissue.

GLONUS TUMOR

deen seated pink or paroxysmal

pu I which case pains in the case pains in the case pains in the case renderness is absent in most of them (Weidman and Wise, Ejster tenderness is absent in most common sites of the solitary lesions are and Montgomery) The most common sites of the solitary lesions are



Fig. 231 Glowius tumor There are numerous vascular lumina of varying are lined by a single layer of endothelial cells and surrounded by proliferating rows of glomus cells. However some of the masses of glomus cells do not show a central vestel (X200)

the nul hed and the fingertups however the lesions may occur else where

Histopathology. The glomus tumor represents a beingn tumor of the currineous glomus a structure composed of an arterial segment the bucquet Hoyer carul and a venous segment. The normal Sucquet Hoyer canal powesses a narrow lumen lined with a single layer of endothelial cells and a tight, number of glomus cells (see page 24). Glomus cells have a faintly cosmophilic cytoplasm and large, oral pale nucles with a distinct thornatin structure, Thus they resemble cpitheloid cells. They are richly supplied withinonmyclinated nerve fibrils.

A glomus tumor is composed of Vascular lumina and numerous

Mesodermal Tumors

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glomus cells As a rule the lumina are small but sometimes especially in those cases with multiple lesions, the lumina are of considerable size (Weidman and Wise, Eyster and Montgomery). The vascular lumina are lined by a single layer of flutened endotibelaticells and usually, by several layers of glomus cells (Fig. 231). Some of the vessels closely resemble the Sucquet Hoyer cruals of the normal glomus. In many areas the glomus cells proliferate arregularly from the viscular walls into the connective tissue stroma of the tumor. In addition makes of glomus cells without central lumina and scattered glomus cells are present in the stroma.

They innective tissue stroma of the tumor is loose, edematous and contains scattered fibroblasts and glomus cells. It may show marked mucoud degeneration. Special straining for herve fibers (Bodian stain) will reveal numerous nerve fibers, most of them nonmyelimited. They show considerable branching and terminate as fine fibrils around the glomus cells.

HEMANGIOPERIC\ TOMA

This rare tumor may arise wherever there are capillaties. Its most common sites are the skin and the subcutaneous tissue. Hemangio

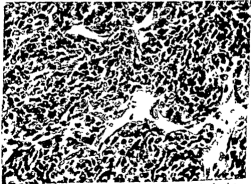


Fig. 232 Hemangiopericytoma. The capillary lumina are lined with a single layer of endothelial cells and are surrounded by irregularly proliferating closely packed pericytes. Most of the pericytes are spindle shaped (2/400).

pericytomas have no diagnostic clinical appearance. They are of varying size firm and nodular Some are benign some are gradually growing and some even metastasize.

Histopathology The tumor is characterized by the presence of vandothelial tubes and sprouts surrounded by triegularly proliferat

In some cases the pericytes show frequent uniton, in our and a vasion into-parentar lumina so that the tumor must be regarded as malignant or at least patentially malignant (Fisher Laulman and Mason). In one instance sudespread metastases resulting in death have been reported (Forrester and Houston).

• ... Pataminat

in the glomus tumor show a <u>more orderly attringeme</u>nt and are more <u>uniform</u> in appearance being round to ovel and never/spindle shaped

KAPOSIS SARCOMA (MULTIPLE IDIOPATHIC HEMORRHAGIC SARCOMA)

kaposis sarcoma consists of an eruption of multiple bluish red of dark brown nodules and plaques. The lesions not infrequently show a verticeous surface. They may undergo affectation. Spontaneous in volution of some of the lesions occurs occasionally. The sites of predilection are the distributions of the extremities but other areas of the Air may also become involved.

Visceral lesions occur in about 10 per cent of the cases (Tedeschi lolsom and Carnicelli). The most frequent sites are in order of fre quency the guttomatismal tract the liver the lungs and the retro peritoneal and the mesenteric lymph nodes (Dorffel). In rare in stances there may be useeral lesions of kapous satromal without utaneous involvement (Tedeschi Folsom and Carnicelli).

Histogeness The Instogeness of Aposts sarcoms is not fully activated Avidely accepted view (with which the author agrees) is that Apost sarcoms as a benien angiomitosis arising from embronal vascular cells and that the lessons are autochtomous in origin rather than metastatic (Gilchrist and Ketron Lang and Hast

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hofer, Becker and Thucher Tedeschi, Folsom and Carnicelli) Dit senting views are held by Vautrier and Diss who regard kaposa's streomy as a neutro of the and related to the lymphoma group of proposed originally by kaposa has been abundoned by most writers



Fig. 235 hapous surrours early granulomatous stage. The capit butes are increased in size and number. Their endothelial cells are large. A diffuse chronic, will unantiony infiltrate is present. In the lower cuttral section of the illustration one can see groups of endothelial, cells, unempting, to form, was, bland, vessels, (§ 200).

Nevertheless some authors (Aegetter and Peale, Sachs Azulay and Convit) still maintain that it represents a sarcoma

Evidence against the concept of Kaposi's surcoma as surcoma and in two of the autochthonous rather than the metastate origin of the lesions are (1) the absence of a primary focus that progressively enlarges, (2) the appearance of widely separated lesions in crops (3) the spontaneous regression of some lesions and (4) the fact that histologic examination may reveal very early stages of development in late appearing lesions. Occasionally, however, a lesion may undergo malignant degeneration and then grow as a true surcoma and cause metastases.

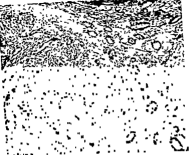


Fig. 234 Kaposi's sarcoma neoplastic stage. On the left, the neoplasta is fibroblastic on the right, it is angiomatous (×100).



Fix 25 Apposts parcoma angiomatous lesson. There are numer our vacular lumina. Most lumina are lined by only a single layer of endothelial cells but some are surrounded also by perithelial cells. The stroma is edematous and contains extravasated erythrocytes (v200).

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hofer Becker and Thatcher Tedeschi Folsom and Carnicelli) Dissenting views are held by Wautrier and Diss who regard kaposa's sarcoma as a neuroascular dysgenesis and by Dorffel who regards it as a disease of the reticulo-endotichal sastem and related to the lymphoma group of diseases. The idea that the disease is a sarcomaproposed originally by kaposi has been alandoned by most writers

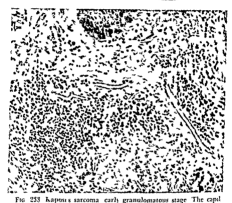


FIG. 233 Kaposis sarcoma early granulomatous stage. The Capin Irries tre interested in size and number. Their endoubtelal cells are large. A diffuse of trone_inflammators infiltrate is present. In the lower central section of the illustration one can see groups of endothelial cells ittempting to form new blood vessels (X*900).

Nevertheless some authors (Aegerter and Peale Sachs Azula) and Convit) still maint in that it represents a sarcoma

Endence against the concept of Kaposis sorroom as sarroom and in favor of the autochthonous rather than the meristate origin of the lesions are (1) the absenct of a primary locus that progressively enlarges (2) the appearance of widely separated lesions in crops (3) the spontuneous regression of some lesions and (4) the fact that histologic examination may receil cert early stages of development in late appearing lesions. Occasionally, however, a lesion may undergo malignant degeneration and then grow as a true streoma and cause metastases.

20 settculum represents young collagen) In old regressing lesions one may find considerable collagemization due to maturation of the young libroblasts. This may lead eventually to fibrosis cicatrization and disappearance of the Ission

If malignant degeneration occurs in a lesion of Kaposi's sarcoma the resulting sarcoma is indistinguishable from fibrosarcomy.

Seven instances are recorded in the Interature in which implooms occurred in patients with Kapou is sarroma. In four of these patients himphops was present (Cole and Crump Hufnagel and Dupon: Sachs and Gray Fischer and Cohen) in two mycoss fun goides (Lane and Greenwood Winer) and in one Hodgkins discase (Greenstein and Conten). There is not sufficient proof that Kapous sarroma and lymphoma are related. It is therefore, best to regard their occurrences.

HEMANGIO ENDOTHELIOMA (HEMANGIOSARCOMA)

Malignant tumors arising from blood vessels are Tare. According to the two types of cells of which capillaries are composed perturbulated is and endothelial cells who types occur hemangio fibrosarcoma and hemangio-modific

Hemangue hibrosarcom's have the histologic appearance of fibro sarcomas with conspicuous capillary proliferation and may there fore be regarded as fibrostromas. The sarcomas which occasionally rise in lesions of kaposis syrcoma are of that type. This leaves themanguo endottleitoma as the only specific malignant tumor arising from blood cisells.

Hemangio dolohehoma usually occurs as a diffusely infiltrating miss which is soft dark red and raised above the surface of the skin ligrous slowly but progressively and may attent large size Metastases or cur.

His is characterized (1) by the principle of the residence of the residenc

for their luming to anastomose (Stom)

The number of sacular lumina varies in different tumors but as a rule sacular lumina are numerous. They are irregular in size and shape Large tortious states may be present (Fig. 236). The vascular channels are funded by large, uspical endothelial cells. In many are is endothelial cells he in several layers and proliferate into the lumina to such a degree that the vascular tubes are completely obscured and cannot be made out as such when routine stams are used. When however, agreed when the sacular and cannot be made out as such when routine stams are used. When however, agreed when same are used.

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Histopathology. The two types of cells of which capillaries are composed participate in the formation of the lessons endothelial and perithelial cells. The perithelial cell is a pericapillary histocyte which may develop into axiitoplast (see page 33). In addition an inflammatory reaction is observed in carly lesions. One magditude the lesions of Kaposi's surcoma into carly granulomatous lesions and Jite neoplastic lesions. The latter may be either 'angiornations, if endothelial cell proliferation predominates, or "fibroblastic" if perithelial cell proliferation predominates.

Invertly granulomatous lesions, the blood lessels of the dermis are dilated and increased in number (Fig. 233). Their endothelial cells are swollen. There is perivascular as well as diffuse cellular infilitation, varying in severity. The inflitrate is composed of lympho cytes, plasma cells and some histocytes and fibroblasts. One may see groups of endothelial cells attempting to form new blood vessels. Frequently, one sees small groups of extravasated erythrocytes and deposits of hemosiderin. The histologic picture in the early stage is not always/diagnostic, however, the presence of extravasted erythrocytes and of hemosiderin in a granulomatous lesion as described above should always make one think of early. Kapos is streoma

In late lesions, the histologic picture may be either angiomatous or fibroblastic, frequently both phases are found intermingled in the same lesion (Fig. 234). In agricumatous lesions one finds numer ons vascular lumina. They vary greatly in size and occasionally are saccular. Most lumina show only a single layer of swollen endothelial cells, but some are surrounded, in addition, by perithelial cells (Fig. 235). The stroma in which the vessels are embedded often is edematous aird usually contains hemorrhages and hemosiderin deposits.

Indibroblastic lesions, one observes marked proliferation of spin dle shaped cells which represent young fibroblasts (Symmers) and are derived from perithelial cells. They he in strands which extend irregularly in all directions. The nuclei viry in size and staining qualities and some of them are atypical. Mirotic figures are present though usually in small number. The histologic picture is thus very much like that of a fibrosircom. One feeture however, distinguishes the fibroblastic lesions of Kaposi's streom; from fibrosircoma and that is the presence of small or large groups of extravisticel cryling cites and of granules of hemosiderin between the fibroblasts. This feature sometimes is not compicuous but a thorough serich or staining for from almost invariably will reveal such areas (see Plate 3). The fibroblasts, being immuture, form only little collagen as a rule but staining with Foot's stain will reveal a rither dense network of reticulum fibers produced by them (Symmers). (As discussed on page

can be seen. In some tumors, one may see endothelial cells extend as invading cords between the fibers of the connective tissue and, occasionally, they may occur in large, solid sheets. The endothelial cells seen in this tumor are large and polyhedral.

The endothelia Meells seen in this tumor are large and payactum and have a well-defined cellular membrane and relatively clear cyto-plasm (Fig. 237). The nuclei tend to be round to oval, pale and vesicular. However, many nuclei are atypical, speing irregular in shape and hyperchromatic. In addition, numerous mitotic figures are usually present. Some of the endothelial cells are multinucleated.

POST MASTECTONY LYMPHANGIO SARCOMA

Lymphangio sarcoma may occur in post mastectomy lymphedema (Stewart and Treves) Several years after radical mastectomy for car cinoma, subcutaneous and cutaneous nodules appear in the edem tous tissue of the arm on the side of the operation. The cutaneous nodules have a bluish color. The chinical resemblance to Kapow's sarcoma may be great. The nodules are not radio sensitive. Metas tasses especially to the lungs, occur.

Histopathology. The nodules are composed of large, atypical cells which, in some pixes form solid problerations and, in others, produce capitlary vessels and lacturity structures. The lumens are usually empty, but, in some cases, contain occasional accumulations of red blood cells. In such cases, the tumor represents a furved lymphangio and, hemangio-endothelioma [Jessuer Z.1 h. and Rein]. Outside the tumor formations, one observes numerous dilated, problerating lymphatics in the dermiss in the subcutaneous rissue and even deeper, in the interminental faster.

Differential Diagnosis. The resemblance to Stemmagio endothe lioma is so great that differentiation is impossible except perhaps on the brus of the amount of red blood cells in the lumina). The tumor differs from kaposis as recoma by the intralymphatic proliferation and greater asypicality of the endotheral cells and by the absence of anymeal filtroblants formations.

6 TUMORS OF MUSCULAR TISSUE

LEIOMYOMA

There are three types of lesomyoma of the skin (1) multiple cu raneous lesomyomas (2) solitary angiolesomyomas and (3) solitary genjial lesomyomis (myomes dartoiques).

Multiple curineous leiomyomas are from pinhead to pea sited, brown or bluish, firm elevated nodules which tend to occur either



The 236 Hemming endothehoma Low magnification. There are numerous viscular humain, on the right side is a large viscular sinus. The vascular channels are lined by large atypical endothelial cells which in some areas proliferate integularly into the luminar (x100).

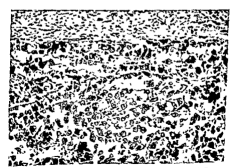


Fig. 237 Hemmigo endotheliona. High mignification of Figure 236 A large anus is shown into which atypical endothelial cells prohierate. Some of the endothelial cells are multinucleated. (X200)

muscle of years and genital leiomyomas from either the muscularis sexualis or the muscularis mamiliae (Stout)

All leiomyomas are composed of interlacing bundles of smooth muscle fibers (Fig 238) Proliferating collagen bundles are often intermingled with the smooth muscle bundles. The muscle fibers composing the bundles are straight or slightly wary and contrin cen trilly located thin very long blunt edged nuclei. The muscle bun



contract is present within a leiomyoma. The muscle bundles of the vein merge with those of the tumor (×200)

finne bine stain may be used Work it . ..

manifulty sew blood vessels

Solurry angioleiomyomas he largely in the subcuraneous tissue, are encapsulated and contain large blood vessels, probably veins, with that muscular walls (Fig 239) Some of these vessels have a stellate lumen because of contraction of the muscular tissue. In some areas, one sees the muscle bundles of the vessels merge with those of the Soluate ---

пофенсара t rather la

438 Mesodermal Tumors

on the back the face or the extensor surfaces of the extremities and usually are arranged in groups. They often are painful and sensitive to pressure



Fig. 238. Lecomorda non-recular type. The tumor is composed of interlicing bundles of smooth muscle fibers. The nuclei of the smooth muscle fibers, re thin doing and blunt edged. Collagen is milter in intermigled with the smooth muscle bundles. Both starn alick vith hemitosylin cosin. To differentiate them, an antine blue stain may be used. (See Plate § 1 (×100).

Solitary angioleiomyomas usually are subcutaneous in location but are adherent to the overlying skin. They rarely grow larger than I cm in drapeter and usually are nontender.

Solutary Schutal leomy omas are located either on the scrotum the hhammapra or rarely on the nipples They may attain considerable size several centimeters in diameter and are nontender

Histopathology Multiple cutaneous leions om is arise from the arectores pilorum muscles solitary angioleiomyomas from the smooth

uenth, shows active downward probleration, even with horn pearl irmation (Fig. 241) (Bloom and Ginifer). This pseudo epithelioma aus hyperplasia has been mistiken for squamous cell carcinoma in veral cases reported in the literature (Eackhoft).



pseudo-epithelumatous hyperplasia. The dermis contains nu merous large pile cells as they are typical of this tumor (×200)

In lesions of the tongue one not infrequently sees are is suggesting of instituous between the tumor cells and mature intuscle fibers (Crane ind Tremblas). This observation has been the reason that granular cell myoblistomas were at first generally regarded as tumors of immature striated muscle cells (myoblasts). However, in recent years, several vuldors have expressed the view that these transitions were suita apprient and that granular-cell myoblationas were composed of eulter shaumania cells or endoncerial fibroblasts and thus were of neural origin (Fust and Custer, Ashburn and Rodger, Bangle).

44n

GRANULAR-CELL MYOBLASTOMA

Granular-cell myoblastomas are solitary tumors occurring most commonly in the tongue, the <u>shin</u> and the subcutaneous tissue (Crane and Tremblay). They usually are benign, but malignant degeneration occurs occasionally (Ross, Miller and Foote). Cutaneous myoblastomas usually consist of a firm, tound, well-crunscribed, non-

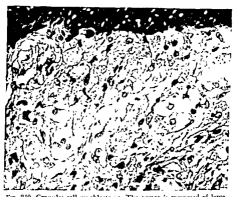


Fig 240. Granular-cell myoblastoma. The tumor is composed of large cells having a pale cytoplasm filled with coarse granules. (X400)

tender nodule, from 0.5 to 2 cm. in diameter, which may lie within the thickness of the skin or may be pedunculated. In some instances, the surface of the tumor is hyperkeratotic Subcutaneous myoblastomas consist of a firm nodule which may or may not be attached to the overlying skin.

Histopathology. On histologic examination, the Kells of the tumor appear large and polyhedral. Most cells measure from 20 to 60 microns in diameter, but some are even larger. They have a pale cytoplasm filled with coarse acidophilic gramules (Fig. 240). The micro are small, round or oval and somewhat vesicular (Cipollaro and Einhorn). Some cells contain more than one nucleus. Delicate strands of collagen surround, the cells of the tumor.

The overlying epidermis usually is hyperplastic and, not infre-



F 6 919 Osteoma cut s Lo v magn ficat on The bone appears lamel



f 13 Osteoma cutis It gh magn ficat on of F gure 21" The bone i ned n many areas by osteoblasts. In the center of the field several os cox lasts in who a niche called Howsh ps lacuna (X°00)

The reasons these authors have given are first, granular cells occa bundles of axis cylinders (Fust and Custer), second groups of groun lar cells have been found within nerve sheaths both within and in the vicinity of the tumor (Fust and Custer, Ashburn and Rodger) and, third, the granules stain slightly positive for lipids, suggesting that they form as a result of the disintegration of axis cylinders and myelin sheaths (Bangle)

In the rare cases of malignant granular cell myoblastomas, one can observe all stages of transition from typical granular cells to malig

observe an stages of transition from typical granular cells to mang nant spindle and grant cells devoid of granules Widespread hema togenous metastases may occur (Ross, Miller and Foote)

Differential Diagnosis. On cursory examination, the large, pile cells of the tumor resemble the foam cells of xanthoma However, the cells of granular cell myoblastoma contain a granular and not a foamy cytoplasm and takefat stains only very faintly Furthermore, in granular cell myoblastoma, the overlying epidermis tends to be hyperplastic rather than atrophic as in vanthoina

7. TUMORS OF OSSEOUS TISSUE

OSTFOMA CUTIS

Cutaneous bone formation may be primary (heterotopic) or sec ondary (metaplastic). Only lesions with primary bone formation should be called osteomas

In primary bone formation bone develops in areas which were not the site of previous lesions. The bone probably develops from em bryonal rests Such lesions therefore represent nevoid tumors or hamartomas (Hopkins, Dietrich Vero, Machacek and Bartlett)

In secondary bone formation, bone develops in areas of tissue de generation Bone may develop in tumors particularly in the calcifying epithelioma of Malherbe (see page 368), in scar tissue (Lifga and Burns) in lesions of scleroderma in various granulomas and in areas of fat nectosis or hemorrhage. Also multiple small foci of ossifica-tion may occur in the skin of the face in prolonged, severe acine with scarring (Leider) In all these instances the bone develops by meta plasia and frequently, though not always calcification precedes the ossification

Osteomas may be single (Dietrich) or multiple (Hopkins, Vero Machacek and Bartlett, Tydens and Rinter) Their clinical appear ance is not uniform Usurilly they occur as small hard plaques or nodules within the dermis or subcutis

Keloid

Garb J and Stone W J Keloids Am J Surg 58 315 1942

Fibrosarcoma

Binkley G W Dermatofibrosarcoma protuberans Arch Dermat & Syph 40 578, 1939

Costa O G Progressive recurrent detmatofibroma (Datier Ferrand) Arch

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Centele II Mahgnant fibroblastic tumors of the skin 'scta dermat venereol 31 suppl 27 1951 (Good review)

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1910
Crill J and Kurma J F. Recklinghausen's disease with unusual symptoms

How

Histopathology On histologic examination, one or several pieces of bone are observed within a lesion. The bone appears limelihed in concentric rings about several Havestain carals. These canals contain blood vessels and connective tissue (Fig. 212). Most of the bone limelihe contain numerous small facture each filled with a cell. These cells called osteocytes possess various shapes, some of them are stellate. Along the margin of the bone one sees many osteoblass which build bone and a few osteoclass which absorb bone. The osteoblasts have small oral or elongated nuclei. As they lay down bone substance they become enclosed in the bone as osteocytes. The osteoclasts have multiple large nuclei and resemble multinucleated foreign body grant cells. Frequently, they lie within deep grooves called Howship's lacume, which extend into the bone substance (Fig. 213).

The tissue surrounding the piece or pieces of bone often is highly viscular and cellular and may contain fat cells so that it resembles bone matrow

Occasionally one may find in osteomas not only bone but also fibrocartilage (Vero-Machacek and Bartlett)

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PLATE 4

Leiomyoma Aniline blue (Vfallory) stain This stain serves to differentiate collagen from smooth muscle. With hematoxylin-cosin both stain red but with aniline blue collagen stains blue and muscle red (×170).





Blue nevus Numerous large, spindle shaped deeply pigmented cells are located in the lower dermis. They are mela noblasts. (×175)

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20

Pigmented Nevi and Malignant Melanoma

Pignuented nevi and malignant melanomas are composed of nevus cells Ventigo, the Mongoliun spot and the blue nevus represent special types of pigniented nevi which will be discussed at the end of this chapter

PIGMENTED NI VIS

From a histologic point of view, it is practical to divide pigmented next into intraderinal (resting) next and junction (active) next. However, intermediary forms so called compound next, are common it is thus understandable that one cannot always predict from the climical appearance whether a nexus is of the intraderinal type or the innertion type.

The sattradermal nevus, as a rule, is a circumscribed, smooth or slightly verviceous elevation which may or may not contain a number of that s and varies in color from normal skin to dark brown Intra dermal nevi are very rare on the springs, the soles and the segundal Nevi located in these locations almost always are junction nevi

The function nevus is a relatively flat or slightly raised, smooth, pigmented mark that is devoid of hairs. Its color is light brown or brownish black rarely slate blue or bluish black. The majority of the marks that are slate blue or bluish black are not junction next blue next.

Histogenesis of the Nevus Cell. In former years, the nevus cell had been thought to be of epidermal origin. Into believed that nevus cells were modified basal cells that had migrated (dropped.od.) from the epidermis into the detimis (Abtropfing theory). Today there are but fen adherents of this theory. (Allen). The great migration in 1926, that both the melanocytes in the epidermis, the so-called clear cells, and the nevus cells are of neural origin (see page 5). Masson stated that nevus cells may develop from two sources from melanocytes in the epidermis and from schwaman cells of cutaneous nerves. He believed that junction nevi develop exclusively from

melanocytes whereas compound and intradermal nevt develop from both melanocytes and schwannian celly. So long as there was ringra tion of melanocytes from the epidermis into the dermis the nevus was a compound neuts but when this migration ceased it became an intradermal nevis. This compound and intradermal nevis according to Masson have a dual origin from two primordia which fused together. This dualistic view of the origin of compound and intra-

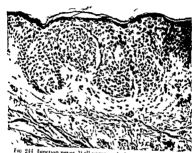


Fig. 244 Junction news Well circumstrated news cell nests are present in the lover epidermis. Otherwise the epidermis appears normal. This type of junction news is not apt to become malignant. (X200)

dermal next has been questioned by some authors (Lund and Stobbe) who believe that all pigmented next develop solely by dropping off

As evidence in favor of his theory of the neural or any

in the deep por iroid tubes and

sons theory is the observation by Berkheiser and Rappoport that of superficial cutaneous nerves cells may be found profiferating within the perineural sheath of superficial cutaneous nerves

20

Pigmented Nevi and Malignant Melanoma

Argumented nevi and malignant melanomas are composed of nevis cells of the Mongolian spot, and the blue nevus represent special types of pigmented nevi which will be discussed at the end of this chapter

PIGMENTED NEVUS

From a histologic point of view, it is practical to divide pigmented nevi into introdermal (resting) nevi and junction (active) nevi. However, intermediary forms so-called compound nevi, are common it is thus understandable that one cannot always predict from the climical appearance whether a nevus is of the intradermal type or the junction type

The satradermal nevus, as a rule, is a circumscribed, smooth or slightly vertucous elevation which may or may not contain a number officiates and varies in color from normal skin to dark brown Invadermal nevi are very rare on the balms the coles and the Jenualia Nevi located in these locations almost always are junction nevi

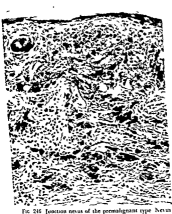
The function nexus is a relatively flat or slightly raised, smooth pigmented mark that is devoid of hairs. Its color is light brown or brownish black, rarely state blue or bluish black. The impority of the marks that are state blue or bluish black are not junction next blue new.

Histogenesis of the Nevus Cell In former years the nevus cell had been thought to be of epidermal origin vann beheved that nevus rells were modified basil cells that had migrated (dropped off)

of investigators have accepted the theory may proper Masson in 1926, that both the melanocytes in the epidermis the so called clear cells, and the nevus cells are of neuril origin (see page 5)

in 1920, that ooth the meanneytes in the epiderinis the colear cells, and the nexus cells are of neural origin (see price 5) Masson stated that nexus cells may develop from two sources from melanocytes in the epidermis and from schnannian cells of curmeous nerves. He believed that junction next develop exclusively from

owined homogeneous cytoplasm. The nucleus is large, the ept ies may



cells he diffusely scattered in the lower epidermis, which appears disorganized No actual invasion of the nevus cells into the dermis is present however. The upper dermis contains a bandlike inflammatory influence intermingled with melanophores (It is possible that this type of junction nevus actually represents the earliest phase of a malignant melanoma) $(\times 200)$

JUNCTION NEVES. This form of pigmented nevus, well described by Fbert by Traub and keil and by Allen and Spitz, is character ned by the active formation of news cells in the lower epidermisie at the epidermal-dermal junction. There are two types of junction ness between which, however, there are transitions. In one type, the newly formed nevus cells are present largely as well-circumscribed 452

Histopathology. In a function (active) nevus, there is active forma tion of nevus cells in the basal layer of the epidermis (i.e., at the epidermal dermal junction) and no nevus cell nests are found in the dermis In a sympound nevus, there also is junction activity, but, in addition, well formed nevus cell nests are present in the dermis. In an intradernial nevus junction, activity is no longer present and the

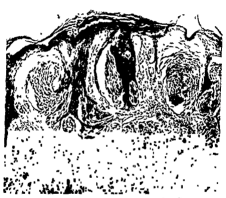


Fig 245 Nevus pigmentosus compound type The cells composing the intra epidermal and the subepidermal nevus cell nests are spindle shaped and resemble schwannian cells. The nevus cell nests suggest Meissner tactile bodies (×100)

nevus cells are all located in the dermis. However, completely intia dermal (resting) nevi are rare if serial sections are made it will be seen that nearly all intradermal nevi show at least a few areas of

junction activity

That the development from junction nevus into 1 sompound nevus and, further, into an untradermal nevus is a matter of aging of a nevus is suggested by the observation of Lund and Stobbe that a much higher percentage of nevi show junctional activity in children than in adults Only nevi located at the distal portions of the extremities were found consistently to be junction nevi, even in adults

The typical flevus cell is oval or cuboidal in shape

suggest Meissner tactile bodies (Fig. 245). Although all junction next have a certain potentiality to become malignant, because they are active next, the danger of malignant degeneration is slight in this type of junction nexus with its regularly formed nests of nexus cells.



110 _10 Netus pigmentosus, intraderinal type. In the upper dermis, the news cells he in nests and cords. In the lower dermis, the news cells are arranged more loosely and embedded in fibrous tassue. (×100)

in the second type of junction nexus, the premalignant junction nexus of Allen and Spitt, in which the nexus cells headiffusely scat tered in the lower epiderius, the lower epiderius appears disorganized by the presence of numerous vacuolated nexus cells with irregularly shaped nuclei (Fig. 246). The nexus cells usually contain a considerable amount of melanti. The border between the epiderius and the derins is somewhat irregular, but no actual invasion of

nevus cell nests within the lower epidermis while in the other type the nevus cells are scattered diffusely through the lower epidermis This latter type of junction nevus has been called premalignant junction nevus by Allen and Spitz

In junction next with will circumscribed nexus cell nests in the lower epidermis the residual epidermis appears essentially normal

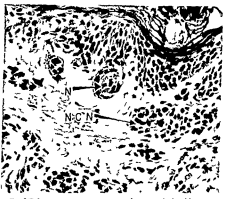


Fig. 247. Nesus pigmentosus compound type. In the basal layer o c sees clear cells (inclinocytes) lying 5 rgly and in nests. Those lying in nests have sliready the appearance of nesus cells. One 1 est (N) is in the stage of dropping off. In all tion typical nesus cell nests. (N C N.) he free in the deemis Considerable amounts of melanin are present in the clear cells and in some of it is nesus cell nests. (X**00)

(Fig 244) Frequently there are nests which he beneath the epider mis but still are in contact with it and thus are in the stage of drop ping off Occisionally in other parts of the same neuts one observes nevus cell nests located free in the dermis representing a development into a compound nexus. The nexus cells contain varying amounts of melaim. Melanophores may be present in the dermis but nownfamiliammatory infiltrate is present. The nexus cells comprising the intra epidermal nests have a regular cuboidal appearance in rare instances however they are spindle shaped and endowed with stroma, so that they resemble schwannian cells and the cell nests

suggest Meissner tactile bodies (Fig. 245). Although all junction next have a certain potentiality to become malignmit, because they are active next the danger of malignmit degeneration is slight in this type of junction nexus with its regularly formed nests of nexus cells



110 410 News pigmentosus intradermal type In the upper dermis the news cells he in nests and cords. In the lower dermis, the news cells he in nests and cords from the lower dermis the news cells are arranged more loosely and embedded in fibrous tissue (X100).

In the second type of junction nevus the premalignant junction nevus of Allen and Spitt in which the nevus cells headingsely scat tered in the lower epidermis the lower epidermis appears disorganized by the presence of numerous variolated nevus cells with irregularly shaped nuclei (fig. 246). The nevus cells usually contain a considerable amount of melanin. The border between the epidermis and the dermis is somewhat irregular, but no actual invasion of

nevus cell nests within the lower epidermis while in the other type the nevus cells are scuttered diffusely through the lower epidermis. This latter type of junction nevus has been called premalignant junction nevus by Allen and Spitz

In junction nevi with Well circumscribed nevus cell nests in the lower epidermis, the residual epidermis appears essentially normal

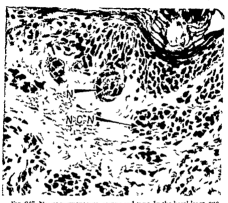


Fig 247 Netus pigmentosus compound type In the basal layer one sees clear cells (melanocytes) lying singly and in nests Those lying in tiests have already the appearance of netus cells One nest (N) is in the stage of dropping of In addition typical netus cell nests (N C N) lie free in the dermis Considerable amounts of melanin are present in the clear cells and in some of the netus cell nests (X200)

(Fig 241) Frequently there are nests which he beneath the epider mis but still are in contact with it and thus are in the stage of drop ping off. Occasionally in other parts of the same news one observes nevus cell nests located free in the dermis representing a development into a compound nexus. The nexus cells contain virying amounts of melanin Melanophores may be present in the dermis but no affirmmentory infiltrate is present. The nexus cells comparising the intra-epidermal nests have a regular cuboidal appearance. In the remainders however, they are spindle shaped and endowed with stroma, so that they resemble schwannin cells and the cell nests

fore can be taken as evidence of the benign nature of the nevus in which they occur. They should not be confused with the giant cells occurring in so-called juvenile melanoma and occusionally, in miliginant melanoma (see page 462). The neutri cells in the Maper por uon of the derius frequently, though not always contain melanin in the presence of much melanin melanophores are often present.



Fig. 250 Vertucous perus pigmentosus. There is licelike downward growth of the epidermis around nests of nevus tells. There are mimerous milimucleated nesus cells. (This is not a sign of malignancy) (x100).

Notaliammytors reaction is present except in the case of mechanical instanton or of secondary infection. In that case, the inflammatory infiltrate however is focal rather than bandliske or diffuse as n is in malignant get-inomy.

In the Word derms, the news cells he farther apart, tend to be youndle shaped and are embedded in fibrous tissue. The fibrous tissue may have the same loose way pale appearance as in neurofibroms if ohen). Within this fibrous tissue, the newsy cells may be arranged in natrion columns so that they suggest setural sheaths (neurous tubes of Masson). In other areas, the fibrous tissue may be in concentre, arrangement (lames foliaces of Masson) resulting in structures resembling Messner teattle bodies (Fig. 249). Monstomery and

nevus cells in the dermis is present. The upper dermis contains numerous melanophores and, in many instances, also in the flammatory infiltrate. This type of junction nevus is often difficult to differentiate from an early malignant melanoma (see page 459) and it possible that it actually represents the earliest phase of a milignant melanoma. In any event, it possesses a definite potentiality to progress into a malignant melanoma.

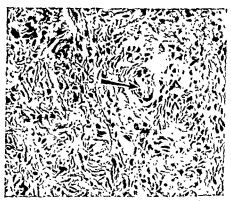


Fig. 249. Nevus pigmentosus, untradermal type. The lower detuns it shown. The nevus cells are lirgely spindle shiped. They are embedded infibrous tissue which has the same loose, with, pile appearance as in neurofibroma. In the center one sees a neuroid structure (N.S.) resembling a Messner trettle book (x.200).

COMPOUND NEWS In a compound news one observes features of both junction news and intradermal news. News cell nests are seen within the epidermis, diopping off from the epidermis as well as in the dermis free from contact with the epidermis (Fig. 247).

INTRADERMAL Nevus Intradermal netishow, in the upper dermis, nests and cords of netus cells surrounded by bindles of collager (Fig. 248). In some instances the netus cells he in dense masses. Not infrequently, one encounters shultinucleated nexis cells showing either clumping or rosette like triangement of small, darkly staining nuclei. These giant cells occur only in well matured next and, there-

melanomas where is initiadermal nevi, which show little or no junction activity, give rise to a malignant melanoma very rively. Especially the so-called premalignant junction nevus which is chiracterized by a diffi-





110 and Margin of an early malignant melanoma. On the left, the epiderms is normal. In the center, because of the presence of many

left beyond the margin of the tumor under the normal epidermis. (x100)

In an and older les but only detrus

dermis

person a numerous at pical necus cells, appears racinolated, tom apart and even disintegrated (Fig. 251). This type of early indiginate melanoma greatly resembles the pre-malignant type of junction necus and differs from it only by the greater at picality of the cells and the beginning invasion into the dermis (see also Differential Diagnosis). As in the premalignant

Kernohan found such structures In 11 per cent of their mitterial of pigmented nest. With special Merie stains such as the Bodian stain one frequently can see nerve fibers in the fibrous tissue. They may occisionally be seen in close relation to the lames foliacies.

An occasional intradermal nexus is devoid of nexus cell nests in the upper dermis and shows only spindle shaped nexus cells embedded in abundant fibrous tissue. Differentiation from a neuro fibroma may then be difficult on a histologic basis (Cohen Becker) Such next, are referred to as neural nexu.

The piderms over intridermal nevi may be normal but often is flutened because of pressure from below. In some nevi the epidermis shows hyperkeritosis pipillomatosis and lacelike downward growth (verticous nevus pigmentosis) (Fig. 250). In others large har follules are present (nevus pigmentosis et pilosis).

MALIGNANT MELANOMA

Miligiant melanoma may arise as such or may develop from a pagmented nexus. If a malignant melanoma arises from this memed nexus such nexus is almost invariably of the <u>minetion</u> type. Chincil evidence that a malignant change is occurring in a pigmented nexus is presented by an increase in the lisize of the lesion an increase in the depth of pigmentation and not infrequently the development of milinflaminatory border with or without spilling of pigment from the lesion into the surrounding skin.

Early malignant melanoma is tharacterized by a gradually enlarging deeply pigmented nodule usually surrounded by crythema Mater the lesion becomes fungating and satellite lesions may appear Ulceration is a late symptom. In occasional instances, hyperpagment

tion is slight or absent

Metastrais takes place at first through the lymphatics resulting in infoliement of the regional lymph nodes. Blood spread is a lite event and may be absent until nearly the end. When it occurs metastrases are usually widespread. The lines the lings the brain and the skin are the most common sites of hematogenous metastrase.

Malignant melanoma is rare before puberty and when it occurs it is usually elimically being n Tor instance in a series of 13 histologically unalignant melanomas in children published by Spitz only one

death occurred

Histopathology The mahgrant changes invariably begin at the definal epiderical junction arrespective of whether a mahgrant mela norm starts as such or develops from a pignicited nexus (Miescher) For this reason pigmented nexis with considerable junction activity like the junction nexus predispose to the development of mahgrant

melanomas, whereas antradermal neva which show little or no junction activity, give rise to a milignant melanoma very rarely. Especially ed by with

a die

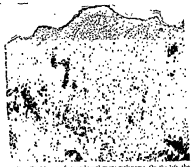


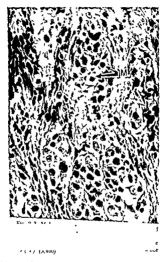
Fig. 251 Margin of an early manginant melanoms. On the left, the epidermis is normal. In the center, because of the presence of many

left beford the margin of the tumor under the normal epidermis (×100)

In an early malignant melanoma and at the advancing border of older lesions one may find considerable changes within the Codermis but only slight invasion of the dermis by union cells. The lower equidermis, because of the presence of numerous asypical nesses cells, appears vacualized, torn apart and even disintegrated (Fig. 251). This type of early malignant melanoma greatly resembles the pre-malignant type of junction nesses and differs from it only by the greater asypicality of the cells and the beginning invasion into the dermis (see also.) Differential Diagnosis.) As in the premalignant



irregularly shiped, often deeply pigmented nests of atypical nesus cells which may show mitotic figures (Fig. 253). The upper epiderinis may be invaded and become so permeated with tumor cells that it



disintegrates and ulceration results (Fig. 254). In addition, there is deep penetration of the dermis by atypical nexus cells. Simultane only with the downward penetration of the nexus cells, the epidermis only the few to by the down.

The size and the shape of the Yumor cells in the dermis show great

460

type of junction nevus, one finds in the upper dermis, close to the epidermis, a dense, bandlike inflammatory infiltrate intermingled with many melanophores. This inflammatory infiltrate often extends for a distance away from the malignant melanoma under the normal



Fig. 252 Malignant melanoma. Low magnification. There is considerable junction activity. One finds not only dropping off of new jeels and news cell nests downward into the dermis but also migration upward into the stratum milipidin. The tumor cells are largely of the cuboidal type and he in alcedra formations. (×100)

epiderinis (l.ig. 251). It is of interest that once deep imasion of the derit.

| Mato | Leells his taken place, the influm and in well progressed malignant there is extension of the tumor.

In an advanced malignant melanoma there is also considerable irregular function activity (Fig. 252). The epidermis comains large,

in malignant melanoma usually is strongly positive in the cells near the dermo-epidermal junction. The tumor cells deep in the dermis, however, react weakly or not at all (Miescher)

Inwhildren malignant melanoma is uncommon and even lesions which histologically have the appearance of malignant melanoma



fusiorm and he in irregular strands. The tumor cells are resembles careed and the control of the

cause metastases only in the rarest instances. These malignant mela nomas generally referred to as <u>juvenile melanomas</u> different in about c

giant cells noma (Sputz, Haber) These giant cells have a round, oval or stellate variation Nevertheless two types of cells can be recognized clearly a cuboidal and a fusiform type. Although most tumors show both cell types almost invariably one type greatly predominates. Predominance of cuboidal cells is much more common than predominance of fusiform cells. The cuboidal cells tend to be in alcohar formations (Figs. 252 and 253). The fusiform cells tend to be in irregular braiching strands (Fig. 255). Tumors in which fusiform cells predominate.



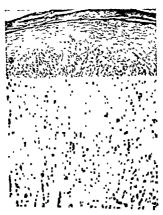
Fig. 254 Malignant melanoma. The epidermis has disintegrated due to permeat in with tumor cells. ($\times 200$)

resemble fibrovircoura but differ from it by the presence of junction activity. Mitoric figures are usually priseant in inalgarant mediatoms but often only in small numbers. They should be searched for be cause their presence is of orest value as evidence that the tumor is mallorant in ismuch as in pigmented nevi mitotic figures are very rare or absent (Mieschet). Busine multimidented grant cells occur occasionally.

The amount of dielaum arises considerably in malignant melanours In some tumors considerable amounts of melanin we present in both tumor cells and inclinophoies. In others particularly in rapidly growing tumors there may be no evidence of melanin in hematoxylin cosin strius. Such tumors have been referred to as amelanotic malignant melanoum. However straining of sections with silver will reveal in most instruces a few cells containing melanin trespective of the amount of melanin present, the doral reaction

in malignant melanoma usually is strongly positive in the cells near the dermo-epidermal junction. The tumor cells deep in the dermis, howevey, react weakly or not at all (Viescher)

Invention malignant melanoma is uncommon and even lesions which histologically have the appearance of malignant melanoma



e u.m. in integular strands. The tumor thus resembles a <u>fibros recomb</u> but differ from it by the presence of junction activity and of considerable amounts of meianin which are located in tumor cells as well as in melanophores (X100).

cause metastases only in the rarest instances. These malignant mela nomas generally referred to as <u>junefule melanomas</u> differ in their histologic aspects from malignant melanoma of additional properties.

variation. Nevertheless, two types of cells can be recognized clearly a cuboidal and a fusiform type Although most tumors show both cell types, almost invariably one type greatly predominates Predomi nance of cuboidal cells is much more common than predominance of fusiform cells. The cuboidal cells tend to be in alreola formations (1 igs 252 and 253) The fusiform cells tend to be in irregular branch ing strands (Fig 255) Tumors in which fusiform cells predominate



Fig 251 Malignant melanomy. The epidermis his disintegrated due to perme ition with tumor cells (X200)

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an execution to the projecting retering each type of the projecting reteringes shows considerable hyperpagmentation and a great increase in the number of clear cells. To junction activity 1 e X-dropping of clear cells is observed. The upperforms often contains melano

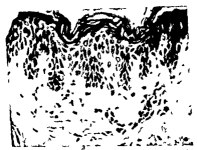


Fig 256 Lentigo The rete dges are elongated Aumerous clear cells (melanocytes) are present in the basal layer (×°00)

plores and sometimes a mild perivascular lymphocytic infiltrate In the case of senile lenugo, the upper dermis also shows basophilic degeneration of the collagen

Whether or not lentigo can progress by the onset of dropping off activity into a junction nexts is not yet fully decided but this appears likely since junction next not infrequently show at the period or the control of the control

MONGOLIAN SPOT

The typical Mongolian spot occurs as a bluish gray discoloration in the <a href="https://link.great.org/link.gr

shape and an acidophilic cytoplasm with one or more large, vacuo lated, irregularly outlined nuclei. They differ from the giant cells of intradermal nevi, which have small, deeply bisophilic nuclei, and from those found occasionally in malignant melanomy, which are larger and more bizarre.

The histologic appearance of metastatic lesions of the skin differs from that of primary lesions by the absence of function activity Metastatic lesions never show any inflammatory infiltrate (Divon)

Differential, Diagnosis. Differentiation of early malignant mela noma from annetion nevus may cause considerable difficulties. That is easily explainable a junction nevus may develop into a malignant melanoma If this development proceeds slowly, as it often does tumors in an intermediate stage of development result. The type of junction nevus most likely to undergo malignant degeneration is the one with diffusely scattered nevus cells in the lower epidermis, the so called premalignant junction nevus (page 455, 1 ig 246) This type of junction nevus, like early malignant melanoma, has numerous vacuolated nevus cells with atypical nuclei in the lower epidermis which thus appears disorganized It may show, in addition, a band like inflammatory infiltrate in the upper dermis. It differs from malignant melanoma by the absence of invasion of the nevus cells into the dermis and by the absence of mitotic figures. However, these are not absolutely reliable criteria, and in some instances a decision as to whether malignancy exists already is impossible. There are some authors, like Miescher, who are inclined to regard as malignant any junction nevus showing a bandlike inflammatory infiltrate. Accord ing to Miescher, this is evidence that aggressive invasion is being fought off It matters little, however, whether such junction nevi are called premalignant or 'early malignant", it is important that they require wide excision

A fighty malign intramelanous melanoma may be difficult to differentiate from a highly malignant fibrosarcoma or a Grade IV smidle cell squamous cell carcinoma. For their differentiation see

under l'ibrosarcoma page 110

LENTIGO

Lentigo is a smooth not infiltrated dark brown mark usually measuring only a few millimeters in drameter A juvenile and a senile type occur fuvenile lentigines begin to appear in childhood and occur on all parts of the body Senile lentigines (three spots) occur in old age on the dorsa of the hands on the forearitis and on the face. Both types are benight Clinical differentiation from junction neuts may be impossible

Histopathology. The histologic findings are the same in Juvenile and enile lenting. The rete ridges appear elongated and chile shaped (Fig. 256). Eccentre thimb like bades may project from them (Peisler and Becker Cauley and Curtus). The first liver of the projecting rete ridges shows considerable hyperpignentiation and a great increase in the number of cleir cells. So junction activity, ie. X droppingoff of clear cells is observed. The upper dermis often contrains melano-

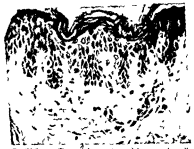


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It hether or not lenting can progress by the onset of dropping off activity into a junction nersus is not yet fully decided but this appears likely since junction ness not interquently show at their periphers, the histologic picture of lentingo

Differential Diagnous Lenngo must be differentiated from ephelis (treckle) Ephelides show hyperpismentation of the basal layer but no elongation of theyeste ridges

MONGOLIAN SPOT

The typical Mongolian spot occurs as a bluish gray discoloration in the <u>lumbovicial region</u>, It is found with great regularity in mem

bers of the Mongolian race but occurs occasionally also in other races It is present at birth and gradually fades

Occasionally, Mongolim spots occur outside the usual location as aberrant Mongolian spots (Cole, Hubler and Lund) They my even be multiple (Carleton and Biggs) or appear during adult life and gradually increase in size (Pariser and Beetman)

Histopathology. The dermis shows, especially in its <u>outdoortion</u> fusiform and stellate cells filled with brown pigment granules. These cells lie widely scattered as well as in small groups. They give a post two dopa_teaction (see pige 12), indicating that they are melanocytes.

(Cole, Hubler and Lund)

Formerly, the cells composing the Mongolian spot were regarded as mesodermal melanocytes, as opposed to the epidermal melanocytes forming pigmented nevi. However, the theory of the neural origin of melanocytes has done away with this artificial division into epidermal and mesodermal melanocytes. The cells of the Mongolian spot (as well as those of the blue nexus-see below) are now regarded as melanocytes which in their embryonal migration from the neural crest to the epiderinis fuled to reach the epiderinis but became ar rested in the dermis (Montgomery)

BLUE NEVUS

The blue nexus is a sharply circumscribed, round or oxal, soft nodule of slate blue or bluish black color which measures, as a rule, only a few millimeters in diameter. There usually is only one lesson, although multiple blue next do occur. In rare instances, large, in filtrated plaques have been encountered (Upshaw, Ghormley and Montgomery) Malignant degeneration of blue next is very tare, but it does occur. (Montgomery, and Kahler. Allen and Spitz) Metastases may develop and cause death.

Histopathology The cells composing blue nevi are of the same type as those composing the Mongolini spot. They are dopi positive melanocytes which became arrested in the dermis during their embryonal migration from the neural crest to the epidermis (Mont gomery). The number of these cells is however, much greater in blue next than in the Mongolini spot.

In the blue nevus grently clongated melanocytes he grouped in irregular misses and bundles in the middle and the lower thirds of the dermis (Fig. 257). Occasionally these cell misses, extend into the subcutaneous layer or close to the epidermis. The epidermis, however, is normal. The fuelanocytes are spindle shaped and have long bipolar, occasionally branching processes (see Plate 4). They he predominantly with their long axis parallel to the epidermis. Most of

them are filled with numerous fine granules of melanin, often so completely that their nucleus cannot be visivilized. In addition, inclain phores (chromatophotes) frequently occur within and beside the misses of melanocytes. The melanophores differ from the melanocytes by being shorter and thicker and by containing coverser granules (Fig. 288). In addition, the melanophores are dopa positive while the melanophores are dopa negative.



in al Blue news Low magnification Numerous spandle shaped metanocytes filled with metanin, he grouped in megular bundles in the middle and the Jouen thards of the dermis and in the subcutaneous En (x50).

Blue nevi may undergo fibrons In that case, thick bundles of collagen separate the melanocytes In occasional instances, the presence of a pigmented nevi said of a blue nevi is in the same lesion has been reported (Montgomer) and Kahlerj

The blue color which the blue nexus presente of a series

manic nuclei), blue nevi in which malignant degeneration has no

be ' ' ' asionally also in other races

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them are filled with numerous fine granules of melanin often so completely that their nucleus cannot be visualized. In addition, ofelano phores (chromatophores) frequently occur within and beside the muses of melanoytes. The melanophores differ from the melano



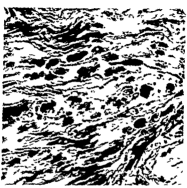
Fig. 257 Rive nevus Los magnification Aumerous spindle shaped melanocytes fulled 1 tilt melanut lie grouped in traggular bundles in the middle and the losser thirths of the dermis and in the subcutaneous fat. (X50)

Blue nevs may undergo hibrosis. In that case thick hundles of collagen separate the melanocites. In occasional instruces the presence of a pigmented nevus and of a blue nevus in the same lesion has been reported (Montgomery and Kabler)

The blue rolor which the blue never presents clinically results from the deep location of the moment have

matic nuclei) blue ness in which mangnant degeneration has oc

curred show areas of necrosis as a diagnostically important feature (Allen). The malignant cells may appear swollen and variobited and may lose their spindled character so that recognition of the lesion as



The 258 Blue nevus High magnification of Figure 2-37 The reason of Figure 2-37 In the melvium granules in addition melanophores filled with Irige irregularly shiped melanin granules in prevent in the middle third of the field whom (x400)

a malignant blue nevus must be based on the presence of residual portions of the original benign blue nevus

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21

Lymphoma and Myelosis

Lymphoma designates a group of malignant tumors arising, usually in multiple foci, from the lymphoid reticular system Alyelosis is a designation for malignant tumors arising, invariably in multiple foci, from the myeloid system

LYMPHOMA

Lymphomas are composed primarily of immature and mature cells of the lymphoid reticular system. The mother cell of the lymphoid reticular system is the lymphoid reticular stem cell. This cell may differentiate into a lymphoid cell (lymphoblast to lymphocyte) or into a reticular cell (reticulum cell to histocyte to fibroblast) (see Chart 2 page 33). The type of lymphoma which results in a given case depends on the degree of immaturity of the cells and on the direction of their differentiation.

The lymphomas may start as a soliting lesion and may remain as such for a long period of time. As a rule, however, the lesions are multiple from the beginning. The multiplicity of the lesions is due to the systemic nature of the disease and usually not to metastasis. Occasionally, however dissemination by metastasis occurs in addition. Acukemia is frequent with some types fare with other types of lymphoma. Leukemia is merely the result of a release of immuture tumor cells into the blood stream and does not represent a separate form of disease. Invariably one finds in patients with leukemia in volvement of the bone marrow. However, marrow involvement may occasionally be found in patients with normal blood pictures (Gall and Mallory).

Just as leukemia is a part of the disease which may be present or absent, so involvement of the Kin is a variable manifestation of lymphonia, found more frequently in some forms of lymphoma than mothers, but a possible occurrence in all In a review of 618 cases of lymphoma, Gall and Mallory found leukemic blood changes in 17 per

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former the tumors are composed entirely of lymphoid reticular cells while in the latter group the tumors show an admixture of inflam matory cells The following classification which is based on that by Mall and Mallory is suggested

Monomorphous group 1 Stem cell lymphoma

2 Reticulum cell lympl oma 3 Lymphoblast c lymphoma

4 Lymphocytic lymphoma

5 Follicular lymphoma (Brill Symmers)

I olymorphous group 6 Hodgkin's disease

7 Mycosis funguides

It is impossible to assign every case of lymphoma to one of these seven types because some tumors are in an intermediary stage of differentiation or show features of more than one type Furthermore different types of lymphoma may be encountered in different areas of the same patient and not infrequently as the disease progresses the lesions may become less differentiated and require reassignment to another type (kenn Gall and Mallory Herbut Miller and Erf)

The terms round-cell sarcoma lymphosarcoma and reticulum-cell sarcoma formerly were and occasionally still are used as designation for single tumors of lymphoma. The use of the term sarcoma is mis leading however because even in the case of a single lesion the potentially systemic nature of the disease may become apparent at any time If he le on

o speci e itiu nonspecific lesions Monspecific

lesions which when found in association with leukemia are often referred to as leukemids may consist of macules papules purpuric legions vesicles bullae eczematous lesions and exfoliative dermatins specific lesions may consist of plaques nodules and tumors, in addition however any of the lesions mentioned as occurring as a non specific reaction may show a specific lymphomatous infiltrate. This holds true especially of eczematous lesions and exfoliative dermatitis Thus it is impossible to predict always from the clinical appearance whether a lesion is nonspecific or specific

Histopathology Histologically specific lesions of lymphoma of the skin show either large masses of lymphoma cells patchy accumula tions of lymphoma cells or 30 inflummatory infiltrate intermixed with lymphoma cells. The simphoma cells vary with the type of lymphoma and may be immature lymphoid reticular cells such as

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Lymphoma and Myelosis

I ymphoma designates a group of mall, man tumors ansulg usually in multiple foci from the lymphoid reticular system Alyclosis is a designation for malignant tumors arising invariably in multiple foci from the myeloid system

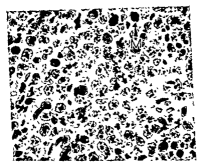
LYMPHOMA

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forms of lymphoma. They occur most commonly in stem-cell lymphoma reticultum-cell lymphoma, lymphoblastic lymphoma and in the tumor stages of Hodgkin a disease and mycosis lungoides. The tumor matter may be sharply demarcated, but usually there are small, outlying islands of tumor cells and, in addition, single rows of lym



Fic. 200 Stemcell lymphoma. High magnification of Figure 239. The tumor cells posses abundant, pale training cytoplasm and large round nucle. The nucle: contain alchest dustike chromatian and one or several promisent nucleoli l'Unione figures are numerous. A large aispical mitotic figure (W) is present in the upper right corner (X400).

phona cells which extend from the main rumor mass no at

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th ... and may occur even in the subcutaneous cissue. They usually show a blood vessel in their center around which the infiltrate lies as a thick sheete.

An inflammatory infiltrate intermixed with tumor cells is the usual picture in Hodgkin's granuloms and in the erythematous and the

stem cells, reticulum cells and lymphoblasts, or mature lymphoid reticular cells, such as histocytes and lymphocytes. Usually attpical cells and a varying number of mitotic figures are present conspecific lesions consist of an infirmmatory infiltrate without immiture cells.

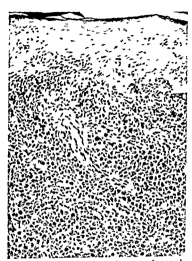
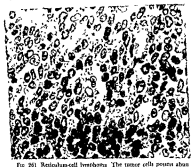


Fig. 259 Stem cell lymphoma. Low magnification: A large mass of loosely packed tumor cells is present in the dermis (×100).

It is probable that most if not all nonspecific lesions result from him phoma cells (Gates). It may be assumed that the himphoma cells either have been overwhelmed and removed by the inflammatory infiltrate which they themselves have provoked or do not differ sufficiently in their appearance from the inflammatory cells to be recognized as hymphoma cells.

Large masses of lymphoma cells in the dermis and the subcutaneous layer, which chancelly appear as cutaneous tumors may occur in all

ture reticular cell (histocyte) [The author prefets to call the immature reticular cell a reticulum cell, and the mature reticular cell a histocyte. However, it should be kept in mind that main authors use the term reticulum cell also for the mature reticular cell (see page 35)] The reticulum cells in reticulum-cell lymphoma have an cosmophilic cytoplasm the border of which tends to be irregular in



rio 201 keticulum-cell lymphoma. The tumor cells possess abun drint pale staining cytoplasm and variously shaped nuclei. Some are round but most are oval or kidney shaped. The nuclei are pale stain ing and possess a distinct nuclear membrane. (X400)

outline suggesting ameboid propensities. The inuclei differ in configuration a leu are round more are oval and still others are kidney shaped. They are pale staining and appear esecular because of the presence of a well defined, strongly basophilic nuclear membrane (rig. 261). They thou a moderately hervy chromatin network, nucleon are rarely cirdent. A moderate number of mutotic figures is present. The reteculum cells, being intimature, form only a scanty reticulum network. In most tumors a fair number of mature reticular cells (histocites) are present. Tumors containing many histocytes may have a well-developed reticulum network. Lymphocytes are sometimes also found presumably evidence of exudative reaction (Wayson and Weithman, Gall and Mallory, Director and Kern.)

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plaque stages of mycosis fungoides. This type of infiltrate, too, tends to have a patchy arrangement

So called nonspecific lesions, which show a nonspecific chronic in flammatory infilitate, occur in the skin, particularly in early mycosis fungoides, in Hodgkin's disease and in lymphocytic lymphomi

I STLY CELL LAMPHOMA

Stem cell lymphoma produces, as a rule, tumorlike lesions In about one third of the cases, it arises as a single lesion, either in the skin or elsewhere, and may remain localized for some time. I eukemin is very rare

Histopathology. The lessons are composed of large masses of cells (Lig 259) The vast majority of cells are stem cells, but most tumors show an admixture of reticulum cells. Stem cells are large in size and possess abundant, pale staining cytoplasm. They may be separate from one another, but frequently the cytoplasm of neighboring cells is fused. Their nuclei are large, from two to four times the size of a normal lymphocyte, round and filled with delicate, dustlike chro matin They usually possess one but occasionally several large, deeply staming nucleoli (I ig 260) Typical and atypical mitotic figures are numerous Stem cells are too immuture to form effectivitium fibers (Gall and Mallory)

RETICULUM CELL LAMPHOMA

Tumorlike lesions are the most common type. In about one third of the cases reticulum cell lymphoma, like stem cell lymphoma starts as a solitary lesion. Occasionally, the first lesion or group of lesions occurs in the skin (Director and Kern)

If leukemia develops in reticulum cell lymphoma, it is of the monocytic variety, since the blood monocyte syderived from the reticular group of cells (Herbit and Miller) True monocytic leu kerner is sometimes referred to as the Schilling type of monocytic leukemia, in contrast to the negeli type of monocytic leukemin m which monocyte like cells arise from myeloblasts and which there fore is a form of myeloid leukemia (Montgomery and Watkins) (see page 195) In the presence of monocytic leukemin the skin frequently shows purpure vesicular and papular lesions (Herbut and Miller) In addition plaques and tumorlike lesions may be present (Wayson and Weidman) Exfoliative dermatitis may occur (Montgomery and Watkins) Swelling and ulceration of the gums are frequent

Histopathology. In tumorlike lesions, large compact masses of cells are present. The type cell is an immatine reticular cell (reticular lum cell) This cell is smaller than a stem cell but larger than a ma

ture reticular cell (histocyte) [The author prefers to call the immature reticular cell a reticulum cell, and the mature reticular cell a histocyte However, it should be kept in mind that many authors use the term reticulum cell also for the mature reticular cell (see page 35)] The reticulum cells in reticulum cell lymphoma have an cosmophilic cytoplasm the border of which tends to be irregular in

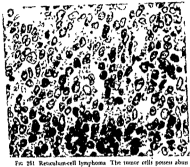


Fig. 261. Reticulum cell lymphoma. The tumor cells possess abut dant prince ound in a not and

outhne suggesting ameboid propensities. The miclei differ in configuration a few are round more are o'vil and still others are kidney shaped. They are palestaining and appear vesicular because of the presence of a well defined, strongly basophilic nuclear membrane (Fig. 261). They show a moderately heavy chromatin network, nu cloth are rarely evident. A moderate number of mitotic figures is present. The resiculum cells, being immature, form only a scanty reticulum network. In most tumors a fair number of mature reticular cells (histocytes) are present. Thimosy containing many histocytes may have a well-developed reticulum network. Lymphocytes are sometimes also found presumably evidence of exudative reaction (Wayson and Westman Gali and Mallory, Director and Kern).

In small cutaneous lesions, as they occur particularly in association with monocytic leukemia, the infiltrate is less extensive and the cells are of smaller size. Nevertheless, the cells have the same appearance as in the large lesions having indented or kidney shaped, pale nuclei with a distinct nuclear membrane (Hubler and Nether

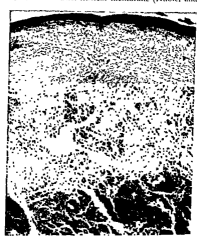


Fig. 262 Lymphoblastic lymphoma. Low magnification. Masses of densely procked ruman tells are present in the lower dermis (X50)

ton). In the presence of monocytic leukemia, extravasation of erythrocytes is frequently found

Differential Diagnosis. It is important not to confuse reticulum-cell lymphoma with reticulo endothehosis (histocytosis) of the Letterer-Siwe and Hand Schuller Christian types (see page 262). Reticulum-cell lymphoma and reticulo endothehosis are occasionally so difficult to differentiate histologically that in the past they have been confused (Swetter, Winer and Cumming). Reticulo endotheliosis, like reticulum cell lymphoma, shows large, irregularly shaped reticular cells (histocytes) but shows no atypical reticular cells, par-

neularly no mitoric figures. The lesions contain as a rule, eosinophils and one may find cholesterol deposits in the reticular cells

3 LAMPHOBLASTIC I YMPHOMA

The cutaneous lesions consist predominantly of nodules, plaques and tumors Purpura is not uncommon

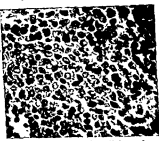


Fig. 263 Lymphoblasue lymphoma. High magnification of Figure 262. The cells have only little cytoplasm. The nuclei lie therefore more densely packed than in stem cell lymphoma.

es procsees a few symphocytes are present (X400)

Leukemia develops in a large percentage of cases—according to Gall and Mailors, in 40 per cent. The type is lymphatic leukemia.

Histopathology The cutaneous lesions show assally large masses of tumor cells and occasionally a patchy infiltrate. Lymphoblasts predommate, but in most lesions stem cells and lymphocytes are present in moderate number. Lymphoblasts possess a narrow basophilic rim of cytoplism and large round or slightly indented nuclei. Because of the relatively small amount of cytoplasm, the nuclei he close to the relatively small amount of cytoplasm, the nuclei he close to the relatively small amount of cytoplasm, the nuclei he close to give much much are larger than those of lymphocytes and more uniform in appearance than those of stem cells or reticultum cells. The citromatin in the nuclei of lymphoblasts is distributed rather evenly and much less clumped than in lymphocytes, giving the nuclei

a vesicular appearance (Fig 263) Nucleoli are observed infrequently Mitotic figures are usually numerous. A fine, evenly distributed reticulum framework is occasionally present (Gall and Millory)

4 LAMPHOCYTIC LAMPHOMA

Cutaneous nodules, pliques and tumors occur l'violitive demnititis is relatively common. In addition, nonspecific lesions, especially

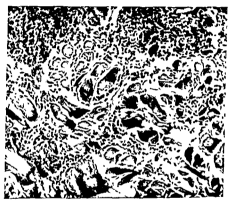


Fig. 264 Lymphocytic lymphoma. In the upper third of the photograph one sees the periphery of a large mass of tumor cells Rows of tumor cells extend from there between the collagen bundles. The cells are indistinguishable from normal lymphocytes (\$200)

papules and petechiae, may be observed. Of interest is the occasional occurrence of generalized herpes zoster in association with lympho cytic lymphoma (see page 491).

Lymphatic leukemia occurs in about half the cases of lymphocytic

lymphoma (Gall and Mallory)

Histopathology. The specific cutaneous lesions may show either large masses of cells (Fig. 264), scattered patches of cells (Fig. 265) or, in the case of exfoliative dermatitis, a diffuse infiltrate in the upper dermis

In lessons showing large masses or scattered patches of cells, the predominating cell is indistinguishable from a normal lymphocyte. Lymphoblasts may be present in small numbers. Minous figures sparse: Thus, the infilirate appears homogeneous. At the periphery of the large masses one frequently sees rows of tumor cells extending between and even around intact collagen bundles in a similar way as it may be observed in scirthous metastatic carcinom.



of lymphocytes are distributed through the dermis. Most patches show a blood vessel in their center (X100)

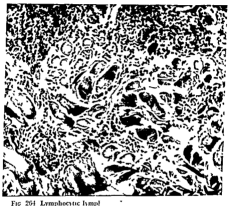
If the infiltrate cobists of scattered patches, some difficulty may arise in differentiating lymphocytic lymphoma from chronic discool lupus erythematosus in which the infiltrate is also patchy. In lymphocytic lymphoma loowever, the patches are distributed indiscriminately throughout the determs without predilection to the upper dermis and the vicinity of the cutaneous appendiages. In addition the patches are uniformly composed of lymphocytes without an admixture of plasma cells and histoocytes as in lupus crythematosus. Furthermore the epidermal changes associated with lupus crythematosus as a bactit.

In instances of exfoliative dermatitis one finds a diffuse infiltrate in the upper dermis. It is composed, in addition to lymphocytes, of

a vesicular appearance (Fig 263) Nucleoli are observed infrequently Mitotic figures are usually numerous A fine, evenly distributed reticulum framework is occasionally present (Gall and Mallory)

4 LYMPHOCYTIC LYMPHOMA

Cutaneous nodules, plaques and tumors occur Exfolutive derma titis is relatively common. In addition, nonspecific lesions, especially



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Histopathology. The specific cutaneous lesions may show either large masses of cells (Fig 264) scattered patches of cells (Fig 265) or, in the case of exfoliative dermatitis a diffuse infiltrate in the upper dermis

6 HODGKINS DISEASE

In most instances. Hodgkin's disease affects primitrily and pre dominantly the lymph nodes. In rare instances, the first lesions are noted in the skin (Reimann, Havens and Herbut).

Cutaneous lessons are observed quite frequently Nonspecific lesions however are more common than specific lesions. Most commonly found are papillar lesions are intensely pruritic. Other lesions are externations patches generalized exfoliative dermatitis nodules or tumors. The latter frequently undergo ulceration (Senear and Caro). Generalized herpes roster occurs occasionally (see page 491).

Histopathology In the specific lesions of Hodgkin's disease two stages can be recognized. Hodgkin's grunuloma and Hodgkin's sar coma Hodgkin's streoma may follow Hodgkin's granuloma, but the disease may show the morphology of Hodgkin's streoma at the onset (Jackson and Patker).

Hodgkin's granulom's composed of a polymorphous infiltrate. The majority of the constituent cells are the usual components of a chronic infiltration infiltrate namely eosinophils neutrophils hyphocytes plasma cells histocytes ind fibroblasts. In addition, and the constituent of the constituent of

from stem cells. They occur as mononucleated and multinucleated cells. The mononucleated Sternberg Reed grant cell possesses a very large irregularly, shaped nucleus. The multinucleated Sternberg Reed grant cell has either a double nucleus (mirror image nucleus) or several nucleu of dissimilar size and shape clustered in the center of the cell. The nuclei of both mononucleated and multinucleated Sternberg Reed cells contain a prominent nucleoliss and heavy clumps of thromatin. Mitotic figures occur with moderate frequency in the stem cells reticulum cells ind Sternberg Reed cells. Hodglan's granu loma has a rendency to focal necrosis and in lesions showing this phenomenon phagocytes are quite numerous. Fosinophili are usually but not invariably present. In some cases, they constitute the predominant cell. Collagen production is the result of the natural evolution of maturing histocytes into fibroblasis. Collagen production is sants in the early phieses and progresses steadily until trood fibrous septa a parate the foci of cellularity into islands (Jackson and Parker).

Hodgkin's streoma differs from Hodgkin's granuloma by the pre ponderince of stem cells resiculum cells and Sternberg Reed cells neutrophils, cosmophils and plasma cells, resulting in an infilmma tory infiltrate of banal appearance. However, immature lymphocites and mitotic figures here and there usually reveal the nature of the process. Sometimes one also finds in the lower cutis variously sized patches composed solely of lymphocytes, further facilitating the diagnosis (Keim).

5 FOLLICULAR LAMPHOMA (BRILL-SAMMERS)

Follicular lymphoma, also called Brill Symmers disease, occurs pre dominantly in the lymph nodes, causing considerable enlargement of the affected lymph nodes. The skin is involved in only rare in stances, showing discrete, firm, brownish to reddish nodules (Gall Morrison and Scott)

Follocular lymphoma of the lymph nodes was described originally as a hyperplasia but is now generally regarded as lymphoma, because it may progress into other forms of lymphoma and is ultimately fatal

Histopathology. The enlarged lymph nodes show replacement of the normal architecture by numerous round, follicle like nodules of varying size. These follicles are composed of densely preked lympho blasts and are surrounded by a thin rim of normal small lympho cytes. Mitotic figures are present within them in small number. Very frequently, the follicles are partially separated from the surrounding stroma by fissures. Although these fissures are an artefact caused by shrinkage of tissue during fivation, they are of considerable aid in the diagnosis of follicular lymphoma (Gall, Morrison and Scott).

The entaneous lesions, in rare instances, show the same follicular pattern as the lymph nodes. Usually however, the infiltrate has the appearance of either lymphostic or lymphoblastic lymphoma.

Differential Diagnosis. Follaular lymphoma of the lymph nodes must be differentiated from reactive hyperplasia such as occurs in dermatopathic lymphadentus (see page 75). It differs from the latter by the presence of mitotic figures, obliteration of the sinuses, absence of inflammatory cells and absence of phagocytosis.

Confusion of follicular lymphoma with derimatopathic lymphade nitis has caused several authors to describe erroneously the occur rence of generalized exfoliative derimatitis in Brill Symmers disease (Combes and Bluefarb Rost). These cases actually were instances of idiopathic generalized exfoliative derimatitis with secondary derimation pathic lymphadenitis.

For differentiation of followin lymphoma from lymphocytoma

cutis, see page 491

6 HODGKINS DISEASE

In most instances Hodgkin's disease affects primarily and pre dominantly the lymph nodes. In rare instances, the first lesions are noted in the skin (Reimann Havens and Herbut)

Cutaneous lesions are observed quite frequently Nonspecific lesions however are more common than specific lesions. Most commonly found are papular lessons which are intensely prurine. Other lessons are ecrematous patiches generalized exfolitute dermatitis nodules or tumors. The latter frequently undergo ulceration (Senear and Caro) Generalized herpes zoster occurs occasionally (see page 4911

Histopathology In the specific lesions of Hodgkin's disease two stages can be recognized Hodgkin's granuloma and Hodgkin's sar coma Hodgkin's sarcoma may follow Hodgkin's granuloma but the disease may show the morphology of Hodgkin's sarcoma at the onset

(Jackson and Parker)

Hodgkin's granuloma is composed of a polymorphous infiltrate The majority of the constituent cells are the usual components of a chronic inflammatory infiltrate namely eosinophils neutrophils himphocytes plasma cells histocytes and fibroblasts. In addition

from stem cells. They occur as mononucleated and multimudeated cells. The mononuclented Sternberg Reed giant cell possesses a very hose irregularly shaped nucleus. The multinucleated Sternberg Reed grant cell has either a double nucleus (mirror image nucleus) or several nuclei of dissimilar size and shape clustered in the center of the cell. The nuclei of both mononucleated and multinucleated Sternberg Reed cells contain a prominent nucleolus and heavy clumps of chromatin. Mitotic figures occur with moderate frequency in the stem cells reticulum cells and Sternberg Reed cells Hodgkin's grante loma has a tendency to focal necrosis and in lesions showing this phenomenon phagocytes are quite numerous Eosinophils are usually but not invariably present. In some cases they constitute the predominant cell Collagen production is the result of the natural evolumon of maturen, histocytes into fibroblasts. Collagen production is scanty in the early phases and progresses steadily until broad fibrous septa separate the foct of cellularity into islands (Jackson and Parker) Hodgkin's sarcoma differs from Hodgkin's granuloma by the pre punderance of stem cells reticulum cells and Sternberg Reed cells

over all other elements comprising the tumor. Multinucleated cells and mitoric figures are numerous Fibrosis is absent or minimal Except for the presence of Sternberg-Reed cells, the histologic pic ture may greatly resemble that of stem-cell or reticulum cell lymphoma (Gall and Mallory)



Fig 266 Hodgkin's disease Low magnification. There are two large masses of tumor cells. In addition, numer ous small collections of tumor cells are present throughout the dermis. (×50)

The nodules and the tumors of the skin show large masses of cells in the dermis and not infrequently also in the subcutaneous far (Fig 266). The histologic picture is more apt to be that of Hodgkin's granuloma than that of Hodgkin's sarcoma and is rarely as typical as in the lymph nodes. The number of Sternberg Reed cells, on the presence of which the diagnosis depends, is often small and fibrosis.

is less pronounced. The presence of atypical reticulum cells with mitotic figures and of chronic influmnation may suggest mycosis fungoides Even then, however, a thorough search of serial sections will usually show a few Sternberg Reed grant cells (Fig. 267). It is important not to confuse clumped reuculum or endothelial cells will multimucleated Sternberg Reed, cells. Resiculum and endothelial



In 267 Hodgkus skiesae High magnification of Figure 266 There is a dense polymorphous inflittate I contains many atypical retordism cells and several muotic figures (M F). A mononucleated Sternberg Reed gant cell (M W) on the right (x400) berg Reed gant cell (M W) on the right (x400).

cells have more regularly shaped paler staining nuclei than Stern berg Reed cells (Miller, Wrong)
In the commission

matit

ease c are also because hieriberg Reed cells frequently are absent. In such instances, the histologic picture is either the same as in mycons fungoides or shows merely a nonspecific chronic inflam matory infiltrate.

The papular lesions show, as a rule, merely a nonspecific infiltrate of lymphocytes and histocytes Occasionally, a few atypical reticulum cells with mitotic figures are present

over all other elements comprising the tumor Multinucleated cells and mitotic figures are numerous. Fibrosis is absent or minimal Except for the presence of Sternberg Reed cells, the histologic picture may greatly resemble that of stem cell or reticulum cell lym phoma (Gall and Mallory)



Fig 266 Hodgkin's disease Low magnification. There are two large masses of tumor cells. In addition numer our small collections of tumor cells are present throughout the dermis (×50).

The nodules and the tumors of the skin show large masses of cells in the dermis and not infrequently also in the subcutaneous far (Fig 266). The histologic picture is more apt to be that of Hodgkin's granulomy than that of Hodgkin's sarcoma and is rarely as typical as in the lymph nodes. The number of Stunberg Reed cells, on the presence of which the diagnosis depends is often small and fibrosis.

attophicans visculate (see page 300) develop either as a precursor or

autopinatis i than it the leasons
as a residum of plaque like leasons
Histogenesis The histogenesis of mycosis fungoides is not solved
fulfy Some authors explain the multiplicity of cell types as due to
continued lymphoid and mycloid stimulation (Reimann Havens and concounce () improve the injective standard of personal Parties and Herbut). A most logical explanation is that first suggested by Fraser (1922–1936). He regarded injects fungoides as a form of reticulum cell lymphoma and believed that all other cells represented merely a defense reaction of the tissue against the tumor cells He pointed out that in the early stage of the disease the number of tumor cells is small and the inflammatory defense reaction pronounced. As the dis reaction slackens until finally in the tumor stage the tumor cells proliferate uninhibitedly Fraser's theory finds support in the fact that also in other malignant diseases such as squamous-cell carci noma and malignant melanoma an inflammatory tissue reaction tends to be present as long as the tumor cells are only moderately malignant but disappears as the malignancy of the tumor cells in creases (see Squamous Cell Carcinomia page 330 and Malignant Vielanoma page 460)

Histopathology The division of mycosis fungoides into three stages as described for the clinical picture pertains also to the his tologic picture

In the first or erythematous stage the histologic picture not in frequently shows merely a banal inflammatory reaction so that a diagnosis of mycosis fungoides may be impossible. In some instances however particularly if several specimens are taken for biopsy and serial sections are made areas may be found in which specific changes (described in the next paragraph) are present. A finding which should always arouse one's suspiction of mycosis lungoides is the presence of patches of cellular infiltrate deeper in the dermis than one would expect to find them in a banal inflammatory infiltrate. It must be emphisized that the decision whether or not early mycosis fungoides exists often is a very difficult one

In the second or plaque stage the histologic picture usually is diagnostic. The following six changes may be present. (1) great multi-plicity of cell types. (2) polymorphism of the bissopration (2). òŧ

ŧ'n de

i icto abscesses

The variety of cells present includes histocytes (= mature reticular cells) reticulum cells (- immature reticular cells) lymphocytes neutrophils eosmophils plasma cells and libroblasts. The number of

7. Mycosis Pungoides

Mycosis fungoides affects primarily and predominantly the skin In the late stage, however, the internal organs may show involvement (Berman; Post and Lincoln)

(Berman; Post and Lincoin)
Several authors (Symmers; Gall and Mallory) have questioned the advisability of regarding mycosis fungoides as an entity. Symmers has called mycosis fungoides a "clinical and pathologic nonevistent," and states that cases reported as mycosis fungoides represent histologically "cither Hodgkin's disease, reticulum cell streoma or lymphosis coma." It is true that some cases which clinically appear as mycosis. fungoides prove, on histologic examination, to be one of the other types of lymphoma, and that cases which at the onset show the his tologic picture of mycosis fungoides later assume the histologic chiracteristics of some other form of lymphoma. Nevertheless inycosis fungoides has a distinct histologic picture which in the majority of cases remains as such throughout the disease and is found also in the internal lesions Transformation of one form of lymphoma into an other occurs not only in mycosis fungoides but also in other forms of lymphoma. This is evidence that all forms of lymphoma are related to each other but it does not obviate any classification

Clinically, as well as histologically, mycosis lungoides can be divided into three stages, the erythematous ("premycotic" or 'pre fungoid') stage, the plaque stage and the tumor stage. All three stages may be present simultaneously and, occasionally tumors develop without the previous presence of erythematous or plaque lesions (mycosis fungoides d'emblee) (Eller and Rein)

In the erythematous stage, the eruption may resemble eczema, psoriasis or parapsoriasis or have the appearance of generalized exfoliative dermatitis Most commonly one observes scattered ery thematous, scaling patches of irregular outline. They may be poorly demarcated or show a fairly sharp border. Since such lesions often resemble eczema closely it is always advisable to consider the possi bility of early mycosi fungoides when dealing with an atypical chronic eczematoid eruption. Similarly, in cases of generalized exfoliative dermatitis of unknown genesis the possibility of mycosis fungoides must always be borne in mind Itching is a prominent symptom in most instances of early inversis fungoides

In the plaque stage cut mate well defined, clevated plaques are seen. They may show central cleving sesulting in ringlike lesions

In the tumor stage, one observes tound or lobulated, raised tumors of bluish to brownish red color. They often undergo ulceration

In rare instances of mycosis fungoides, lesions of poikiloderma

vessel in their center. If the patches are fairly large in size, they are strong evidence for lymphoma. An almost pithognomonic finding occasionally encountered is the presence of so-called Partier * micro-abscesses* in the malpighian layer. They consist of small accumula



Fig. 290 Mycous fungoider plaque stage. High magnification of Figure 268. The infiltrate in the upper dermis shows marked multiplicity of cell tipes and polymorphism of the histocytes Atypical reacculum cells with large hyperchromatic nucles (m) coss cells 3 are numerous. There are several Pautier microalscesses in the epidermis (x400).

tions of cells mainly lymphocytes and histocytes (Fig 269) [It should be recalled that the Munro micro-abscesses of psoriasis are located in the horny layer and are composed of neutrophils) The epidermis in the plaque stage usually shows acanthous with elongation of the rete ridges and may have an appearance similar to that found in psoriasis

In the third or tumor stage the infiltrate consists of large masses of cells and may occupy large areas of the dermis and even penetrate

eosinophils varies but often is considerable. Histocytes are numer ous Staining for reticulum with Foot's stain will show, commensurate with the rather large number of histocytes, a well-developed network of reticulum fibers. The nuclei of the histocytes vary greatly in size and shape. In addition, their nuclei may show pyknosis (shrinking), karyortheyis (breaking up into particles, "nuclear dust").

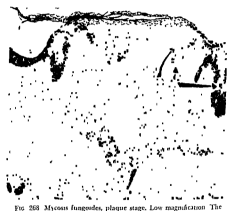


Fig. 208 Mycoss tangoides, plaque stage. Low magnification The infiltrate in the upper dermis is diffuse, while in the lower dermis it consists of variously sized sharply demarcated patches. The epidermis contains several Pautier micro abscesses. (P.A.) (X100)

and clumping Immature and atypical reticulum cells, called mixosis cells, are present. Mixosis cells differ from historicites by having larger, more irregularly shaped and more deeply staining nuclei. Mitotic figures, though not numerous, can, as a rule, be found. (An occasional mitotic figure may occur in a brial inflammatory infilitate, so that the finding of one, two or even three mitotic figures in a section does not necessarily mean mixosis fungioides.) The infilitate in the plaque stage of mixosis fungioides is located largely in the upper dermis just as in a nonspecific chronic inflammation. Fix quently, however, one finds, in addition patches of cellular infilitrate in the lower dermis (Fig. 268). These patches usually have a blood

vesel in their center. If the patches are fairly large in size, they are strong evidence for lymphoma. An almost pathognomonic finding, occasionally encountered, is the presence of so-called Pautrier "mixto absress: in the malpighan layer. They consist of small accumula



of the histocytes. Atypical reticulum cells with large, hyper chromatic nuclei (mycosis cells) are numerous. There are several Pautrier "micro-abscesses in the epidermis (x400)

tions of cells, mainly lymphocytes and histocytes (Fig 269) (It should be recalled that the Munro micro-abscesses of psoriasis are located in the horny layer and are composed of neutrophils) The epidermis in the plaque stage usually shows acanthosis with elongation of the rete ridges and may have an appearance similar to that found in psoriasis

In the third or tumor stage, the infiltrate consists of large masses of cells and may occupy large areas of the dermis and even penetrate

into the subcutaneous layer. The pressure of the infiltrate may de stroy the epidermis so that ulceration results. The infiltrate still shows as a rule, the same characteristics as in the plaque stage except that the number of mycosis cells and of mitotic figures is larger. In some cases, however, the mycosis cells attain considerable size and may possess more than one nucleus so that they resemble mononu.

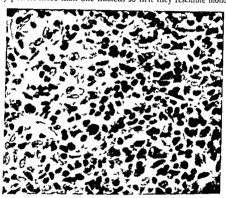
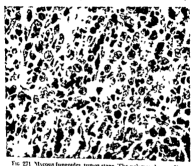


Fig. 270 Mycoss fungoides tumor stage. The infiltrate is polymor phous. There are numerous mycoss cells. Some of them are of considerable size and one located in the center has two in cle. These large mycosis cells resemble mononucleated or multinucleated Stern berg Reed cells. (x400)

cleated or multinucleated Sternberg Reed grant cells (Fig. 270). If such cells are conspicuous the infiltrate may be identical with that of Hodgkin's disease (Friser and Schwartz. Wile and Stiles). In other cases large immature reticulum cells like those seen in reticulum cell lymphoma may be present within the multicellular infiltrate (Fig. 271). Occasionally, immature reticulum cells are so numerous and inflammatory cells so few that the infiltrate is identical with that of reticulum cell lymphoma (Traser and Schwartz).

The histologic appearance of generalized exfoliritive derinatitis in mycosis fungoides may be that of either the erythematous or the plaque stage. If it is that of the erythematous stage a diagnosis of mycosis fungoides may be impossible (Montgomery)

In poikiloderma atrophicans vasculare due to mycosis fungoides, the epidermis is atrophic and shows vacuolization of the cells of the basal layer. A dense infilirate lies in the upper dermis in close approximation to the epidermis and invides the epidermis in some areas. The infiltrate may be that of either the erythemitous or the plaque stage. As a rule sufficient hypical cells are present to permit the diagnosis of mycosis fungoides (Oliver Hazel, Dostrosk), and



Fic 271 Mycosis fungordes tumor stage The polymorphous infiltrate contains many large immature reticulum cells like those seen in reticulum-cell lymphoma (X400)

Sagher) (For a discussion of poikiloderma atrophicans vasculare see page 305)

Involvement of Internal Organs The incidence of specific visceral lesions in mycosis fungoides is difficult to estimate from the interature. Many reviews set the figure too high largely because the tendency exists in the dermatologic literature to regard a case of symphoma as mycosis fungoides just because the primary or predominant lesions of the lymphoma are located on the skin. It should be stressed that for a diagnosis of mycosis fungoides. It is necessary that the majority of the cutaneous leuons either die. were not mycosis fungoides, in a review of autopsies recorded in the literature found visceral involvement in 51 out of 54 cases. Crules, Curtis and Leach, in their own material, found visceral lesions in 8 out of 10 cases of supposed mycosis fungoides, but only 1 case showed the multiplicity of cell types necessary for a drignosis of mycosis fungoides and that case had no visceral lesions. It may be estimated that visceral lesions are found on autopsy in only about 20 per cent of the cases of mycosis fungoides, and usually they are not prominent or widespread.

When involvement of the internal organs is present in mycosis

cutaneous lesions (Ormsby and Timerud) On the other hand, the visceral lesions, just as some of the cutaneous lesions, may be composed largely of immuture reticulum cells and be indistinguishable from reticulum cell lymphoma (Fraser and Schwartz) or may present, due to the presence of Sternberg Reed grant cells the picture of Hodgkm's disease (Fraser, Wile and Stiles, Poulsen)

Of the various internal organs, the subcutaneous lymph nodes are affected most frequently. However, the infiltrate in these lymph nodes is usually not that of mycosis fungoides but is nonspecific in flammatory, showing the histologic characteristics of derimopithic lymphadenitis (see page 75). Nevertheless, the subcutaneous lymph nodes as well as the internal lymph nodes, the spleen the fungs, the liver, the kidness, the gastro intestinal tract and many other organs may be involved by mycosis fungoides (Berman, Gaies, Post and Lincoln). Involvement of the bone marrow however, his never been reported in true cases of mycosis fungoides (Poulsen). For this reason, mycosis fungoides shows no significant hemocytologic changes except that occasionally immature monocytes or lymphocytes are found in the blood. In rare instances, true lymphatic leukemia decelops in the terminal phase of the disease (Lane and Greenwood).

GENERALIZED EXPOLIATIVE DERMATITIS IN LAMPHOMA

Generalized exfoliative derinativis is a fairly common occurrence in lymphoma. It occurs most commonly in lymphocytic lymphoma. Hodgkin's disease and mycosis fungoides but also in reacultum cell lymphoma, particularly when accompanied by monocytic leakening. Therefore, every case of persistent generalized exfoliative determities requires intestigation into the possibility of lymphoma According to Montgomery, 25 per cent of all cases of generalized exfoliative determities prove to be associated with lymphoma. The histologic picture of generalized exfoliative derinatius in lymphoma may be,

in the early stage that of a nonspecific chronic inflammation. In cases in which it is doubtful whether or not lymphoma exists thorough hemosytologic studies lymph node biopsy and sternal marrow biopsy are indicated and additional skin biopsies should be performed at internals.

HERPES FOSTER IN LYMPHOMA

The relatively frequent association of herpes zoster particularly of herpes zoster generalisative with lymphoma is of interest. Herpes zoster is most likely to occur in Hodgkin a disease and in lymphocytic lymphome especially when lymphomic leukemia is present.

hymphoma especially when hymphatta leukemia is present. Histopathology: Histologic examination of the herpes zoster lesons maj show the presence of a hymphomatous infiltrate (Barney Barton and O Leary). In addition autopsy may reveal lymphomatous cells in the intercostal nerve. in the spinal ganghon in the nerve orosts or in the cord segment corresponding to the size of the herpes zoster (Bluefarb). It may be assumed that the lymphomatous infiltrate produces a locus minoris resistentive where the zoster virus focalizes and multiplies.

I YMPHOCYTOMA CUTIS (LYMPHADENOSIS BENIGNA CUTIS SPIEGLER FENDT SARCOID)

This condition occurs in two types: a localized type (Loveman and Fluegelman. Mopper and Rogin) and a disseminated type (Bafver stedi). In the localized type there is either a solitary nodule or a group of nodules in the disseminate type the lesions are scattered widely. The lace and the ear lobes are the sures of prediction. The nodules are soft asymptomatic and radiosensitive Development into a lymphoma does not seem to occur.

Histopathology A hervy infiltrate is present in the dermis usually separated from the epidermis by a narrow zone of normal collagen. The infiltrate occasionally consists only of mature lymphocytes but it most caves there are also restrubing cells. The two types of cells the either intermingled with one another or in a follocular arrange men. In the latter type of arrangement lymphocytes surround islands of reticulum cells resulting in situatures resembling the foliocies of lymph nodes (Fig. 272). The lymphocytes have small round deeply saining mucle lying closely packed because lymphocytes possess only little cytoplasm. The reticulum cells have large irregularly shaped pide nuclei lying in loose arrangement have.

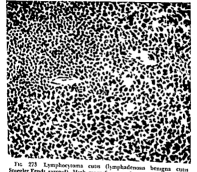
The nature of lymphocytoma cutis is not known. Although it sug gests a lymphoma in its histologic architecture and is highly radio sensitive, it apparently is not a form of lymphoma European authors (Hallam and Vickers; Hellier, Balverstedt) have suggested that it represents a hyperplasia of pre existing rudimentary lymphoid



Fig. 272. Lymphocytoma cutts (lymphadenosis benigna cutis, Spiegler Fendt sarcoid). Low magnification: The infiltrate is composed of two types of cells lymphocytes, which he in the dark struning areas, and retriculum cells, which he in the light staining areas. At the right (LT), the arrangement of these cells resembles that encountered in a lymph follicle (×50)

Differential Diagnosis. The diagnosis is easily established in those cases of lymphocytoma cutts in which the infiltrate shows lymphocytes and retriculum cells in follicular arrangement. Even in cases in which the two cell types he intermingled, the diagnosis usually can be made because the cells composing lymphocytoma, in contrast with those of retriculum cell lymphoma, are mature. Nevertheless, mistakes in diagnosis occur (Director and Kern). The greatest difficulties in diagnosis occur in those cases with a purely lymphocytic infiltrate because it may be impossible to rule out lymphocytic lymphoma. It is thus advisable always to make a diagnosis of lymphocytic und cutts with reservations, except in cases with obvious follicular formations (Loveman and Fliegelman).

Differentiation from follicular lymphoma (Brill Symmers disease) with which lymphocytoma cutts is not related is not difficult since in follicular lymphoma the follicular senters are larger more numerous and outlined more distinctly. Also they are partially separated from the surrounding stroma by fissures which probably are the result of shrinkage of tissue during fixation. Such fissures are not apt to occur in lymphocytoma cutts.



Spiegler Fends sarcoid) High magnification of Figure 272 Lympho cytes lie in the left upper half reticulum cells in the right lower half of the illustration (X200)

MYELOSIS

Myelosis is nearly always associated with leukemia in contrast with lymphoma in which leukemia is often absent. The type of leukemia is myeloid leukemia. On the other hand: cutaneous lesions are less common in myelosis than in lymphoma. Specific and nonspecific cutaneous lesions may occur however. Just as in lymphoma there is no clearcial distinction between the two. The two conditions merge im percepubly and it is probable that most if not all of the nonspecific cutaneous manifestitions result from tumor cells (Gates) (see page 472).

Clinically, the specific cutaneous lesions consist of from pinhead to walnut sized, firm nodules, which may coalesce into pliques and occasionally ulcerate. The nonspecific cutaneous lesions (leukemids) may consist of macules, papules, pustules and purpure lesions

Histopathology. Histologic examination of specific lesions shows the presence of dense misses of cells in the dermis and, occasionally, in the subcutaneous layer. Frequently, rows of cells extend from these masses into the surrounding tissue. In addition, smill groups of cells may be found scattered in the tissue spaces. The majority of cells belong to the myeloid series, but lymphocytes and phagocyte histocytes not infrequently are observed (Ketron and Gay). Some of the cells of the myeloid series are muture, but most are immuture, being either myeloblasts or myelocytes. They may be so immuture that it is impossible to decide whether they belong to the myeloid or to the lymphoid reticular series of cells.

Myeloblasts are about double the stre of a mature polymorphona clear leukocyte. They possess a large, onal or round, pile straining nucleus and relatively little, nongranular cytoplasm. They resemble lymphoblasts, although, as a rule, lymphoblasts fince a conser chromatin structure and less cytoplasm (Nekam). Myelocytes are about the same size as myeloblasts or slightly smaller. They fince, however, a smiller indented or lobate nucleus and abundant cytoplasm con taining granules which may be either fine neutrophihe or coasse cosmophilic. Myeloblasts may or may not give a positive peroxidise reaction (also called oxidase reaction), while myelocytes always give a positive peroxidise reaction.

The peroxidase reaction indicates the presence of the enzyme peroxidase, which occurs in mature and partly matured cells of the myeloid series but is absent in very immuture cells of the myeloid series and in all cells of the lymphoid reticular series. The peroxidase reaction is of great diagnostic value since it is positive in the majority of cases with myelois and always negative in lymphonia. Even very immature tumors of myeloisi issually show at lens a few areas in which the reaction is positive. For the peroxidase reaction, the specimen should be fixed in a 10 per cent formalm solution. Frozen sections are used for staining. On struning peroxidase positive cells show numerous black granules (Ketton and Gay)

Ocasionally, the myeloid cells found in the curineous lesions are more immature than those in the bone mirrow and the circulating blood. This may be regarded as evidence that the curineous lesions in myelosis are autochthonous (Paul and I imazzi). It is probable that the myeloid cells form by myeloid transmutation of local reticular cells (Heller, Lewisolin and Palin).

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CHLOROMA represents a rate form of myelosis in which the myelo blasts and the myelocytes show an unusual tendency to tumor formation. The tumors are found most commonly in the bones especially the flat bones but may occur also in the skin. The peroxidase reaction is usually positive in the tumors. The nature of the green pigment present in the tumors is not yet fully known. It represents an intermediary product in the breakdown of hemoglobin to bilirubin (Goodman and Iverson).

MONOCITIC LEUNEMA OF THE NAEGELITY OF is a form of myeloid leukema in contrast with monocytue leukemia of the Schilling type which is true monocytue leukemia for the Naegeli type the infiltrate in the cutaneous lesions is composed mainly of monocyte like cells so that differentiation from true (Schilling 3) monocytic leukemia in the possible in the blood however myeloiblasts and myelocytes are present in addition to monocyte like cells and the bone marrow shot is myeloiblastic hyperplasia without changes in the reticular cells ofto simple myeloiblastic hyperplasia without changes in the reticular cells (Watkins and Hall). While some regard monocyte leukemia of the Naegeli type as myeloid leukemia with a predominance of monocytes most authors believe that the monocyte like cells differentiate from myeloiblasts. Montgomety and Watkins favor the latter view since they observed cells intermediate between myeloiblasts and the monocyte like cells.

EOMNOPHILIC LEUKEMIA a rare form of myeloid leukemia occa sionally has cutaneous lessons. The cellular infiltrate present in the dermis may consist largely of immature primitive cells with but a few myelocytes and no eosinophilis (Carmel Minno and Cook) or it may contain numerous cosinophilis (Deme)

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Glossary

Cheantholysis Loss of coherence between epidermal cells due to de generation of the intercellular/bridges. It lends to the formation of clefts vesicles and builde within the epidermis. Occurs and the control of the c

(Acant

Alteration cavitaire See Intracellular Edema

Anaplasia Atypical de differentiation of cells occurring in malignant diseases. Anaplastic cells have large hyperchromatic arregularly shaped nuclei and frequently show atypical mitotic figures.

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tion occurs in virus vesicles and is diagnostic of them. See also Reticular Degeneration of Epidermal Cells

Bulla A cavity forming either within or beneath the epidermis and filled with lymph fluid. A Small bulla generally is called a vesicle and a effilike bulla as seen in Darier's disease and senile keratosis a lacuna. About the different modes of formation of bullae see the introduction to Chapter 7.

Cascation necrosis: A type of tissue death in which the affected area has lost its structural outline and consists of pale cosmophilic amorphous finely granular material. Unless the necrosis is far advanced some pyknotic nuclei are still present. Yo in vasion of neutrophils occurs: Cascation necrosis occurs especially in tuberculosis syphilis granuloma annulare and beryl lium granuloma.

Diskerators Faulty, keratinization of individual epidermal cells

by There are two types one occurrying in beingin diseases and the
other in inalignant diseases Meningin diskerators occurs in
Dividual theorem of the properties of the formation of corps ronds and
grains (see page 49) Malignant diskerators occurs especially in Bowen's disease but also in squamous cell carcinoma
and senile kerators and consists of
keratinization of individual cells

for the properties of the p



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(Meantholysis Loss of coherence between epidermal cells due to de generation of the intercellular/bridges. It leads to the forma tion of clefts vesicles and bullde within the epidermis Occurs impemphigus Darier's disease familial benign chronic pem phigus virus bullae and semiler Leratosis, O Acanthosis Increase in thickness of the stratum malpighu

Alteration carnaire See Intricellular Edema

Anaplasia Atspical de-differentiation of cells occurring in malignant diseases Anaplastic cells have large hyperchromatic irregu larly shaped nuclei and frequently show mypical muotic

(a) Ballooning degeneration A type of degeneration of epidermal cells causing marked swelling with loss of the intercellular prindges Acantholysis results and a bulla forms. Ballooning degenera tion occurs in virus vesicles and is diagnostic of them See also

Rencular Degeneration of Epidermal Cells

Bulla A cavity forming either within or beneath the epidermis and filled with lymph fluid A small bulla generally is called a vesicle and a slitlike bulla as seen in Darier's disease and senile keratosis a lacuna. About the different modes of forma tion of bullae see the introduction to Chapter 7

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hum granuloma

Objekeratosis. Faulty keratimization of individual epidermal cells There are two types one occurring in benign diseases and the other in malignant diseases VBenign diskeratosis occurs in Darier's disease and occasionally in familial benign chronic pemplugus and consists of the formation of corps rouds and grains (see page 49) Malignant dyskeratosis occurs espe cially in Bowen's disease but also in squamous-cell carcinoma and semile keratosis and consists of premature and atypical keratinization of individual cells

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Hydropic degeneration of basal cells: See "Liquefaction Degenera tion of Basal Cells"

Hyperkeratosis: Excessive thickness of the horny layer. If caused by excessive formation of keratin, the granular layer is also thickened, as in lichen planus and lupus erythematosus II caused by vetention of the horny layer, as in ichthrosis vul garis, the granular layer is even smaller than normal

Incontinence of pigment: Loss of melanın from the cells of the basal layer due to damage to these cells with accumulation of the melanin in the upper dermis, inside as well as outside of melanophores. It occurs in an idiopathic form in the disease called incontinentia pigmenti, and in a symptomatic form in lichen planus, lupus erythematosus, poikiloderma atrophicans vasculare, Riehl's melanosis and melanodermatitis toxica (Intercellular edema (spongiosis): I'dema between squamous cells

It occurs frequently in inflammatory processes of the skin especially in dermatitis eczema. It does not cause formation of bullae or vesicles but contributes to their increase in size See also Intracellular Edema AIntracellular edema (altération cavitaire). Edema within squamous

causing an increase in the width of the spaces separating them

cells. If severe, it results in reticular degeneration (see below) of the affected cells and in the formation of multilocular hullae Karyorrhexis: Fragmentation of nuclei

Liquefaction degeneration of basal cells A type of degeneration causing vacuolization and disintegration of basal cells. It oc curs in incontinentia pigmenti, lichen planus, lupus erythe matosus, poikiloderma atrophicans vasculare, Riehl's mela nosis, melanodermatitis toxica and lichen sclerosiis et atroph was with its variants kraurosis vulvae and balanitis xerotici obliterans. In several of these diseases, the liquefaction de generation may cause incontinentia pigmenti (see above) In lichen planus lupus crythematosus and lichen sclerosus et atrophicus it may cause the formation of subepidermal bullae (see under Formation of Bullie introduction to Chapter 7)

Metachromasia. The phenomenon of reacting with a different color than that of the dye used for the straining Important exam ples of metachromasia occur in the staining of the granules of mist cells (see 'Urticaria Pigmentosa) and in the staining of mucin (see 'Myxedema) of amyloid (see "Amyloidosis) and of the fibrinoid material in fibrinoid degeneration (see "Acute Systemic Lupus Erythematosus) All four materials

stain purple with toluidine blue methylene blue thionine and cresyl violet Incidentally all four materials because of the presence of polysaccharides stain deeply red with the periodic acid Schiff reaction

Metaplasia Change of one type of tissue into another as it occurs for instance in the formation of bone in scars and in calcifying epithelioma (see Osteoma)

Micro-abscesses Small accumulations of cells in the epidermis/Two types of micro abscesses occur the inicro abscess of lunro goides For their description see those two diseases

Papilloma A tumor or tumor like proliferation of the skin charac terized by papillomatosis (see below) and hyperkeratosis. Five diseases show this type of proliferation nevus verrucosus (Jadassohn) keratosis senilis basal cell papilloma verruca vulgaris and acanthosis nigricans. In typical instances histo logic differentiation of these five diseases is easy but occasion

ally no more specific diagnosis than papilloma can be made Papillomatosis Upward proliferation of papillae causing the surface of the epidermis to show irregular undulation

(Parakeratosis Imperfec nuclei in the horr

is absent in areas Pyknosis Shrinking of nuclei

Reticular degeneration of epidermal cells A process in which severe intracellular edema causes bursting of epidermal cells and formation of smultilocular bulla. The septa inside the bulla are formed by resisting cell walls Reticular degeneration plays a role in the formation of virus vesicles and those of contact dermatitis

@Spongiform pustule of kogoj A multilocular pustule located in the upper stratum malpighii and characterized by the presence of neutrophils inside of edematous epiderinal cells. The cel lular walls of these epidermal cells traverse the puspale like the network of a sponge This type of pustule has a Miagnostic appearance and occurs in acrodermatitis continua of Hallo peau impetigo herpetiformis keratosis blennorrhagica and Reuter's disease Spongrosis See Intercellular Edema

lesions that clinically are hardly visible

c) esicle \ small bulla (see Bulla) No sharp borderline can be drawn between a vesicle and a bulla Generally the term bulla is preferred in histopathology except for very small 500 Glossary

Hydropic degeneration of basal cells: See "Liquefaction Degenera

tion of Basal Cells."

Hyperkeratosis: Excessive thickness of the horn; layer. If caused by excessive formation of veratin, the granular layer is also thickened, as in lichen planus and lupus erythematosus If caused by vetention of the horn; layer, as in ichthysis vul

garis, the granular laver is even smaller than normal Incontinence of pigment: Loss of melanin from the cells of the basal layer due to damige to these cells with accumulation of the melanin in the upper dermis, inside as well as outside of melanophores. It occurs in an idiopathic form in the disease called incontinentia pigmenti, and in a symptomatic form in lichen planus, lupus ery thematosus, poil-floderma atrophicus vasculare, Riehl's melanosis and melanodermatitis toxica.

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Intracellular edema (alteration cavitaire) Edema within squamous cells If severe it results in reticular degeneration (see below) of the affected cells and in the formation of multilocular bullae

Karvorthexis: Fragmentation of nuclei

bullae
Karyorrhexis: Fragmentation of nuclei
Liquefaction degeneration of basal cells. A type of degeneration
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curs in incontinentia pigmenti, lichen planus, lupus erythe
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nosis, melanodermatitis toxica and lichen sclerosus et atroph
icus with its variants kraurosis vulvae and balanitis serotici
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lichen planus, lupus erythematosus, and lichen sclerosus et
atrophicus it may cause the formation of subepidermal bullae
(see under Tormation of Bullae, introduction to Chapter 7)
Metachromasia. The phenomenon of reacting with a different color

atrophicus it may cause the formation of subepidernal bullae (see under Formation of Bullae, introduction to Chapter 7)

Metachromasia The phenomenon of reacting with a different color than that of the dye used for the staining Important exim ples of metachromasia occur in the staining of the granules of mast cells (see Urticaria Pigmentosa) and in the staining of mucin (see Myxedema) of amyloid (see 'Amyloidosis') and of the fibrinoid material in fibrinoid degeneration (see "Acute Systemic Lupus Frythematosus). All four materials

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Villi Elongated and often tortuous papillie which are covered, as a rule, with but one or two layers of epiderinal cells and extend into a vesicle a bulla or a cystic cavity. Formation of villi is observed in Darier's disease, familial benign chronic pemphigus, pemphigus vulgaris, pemphigus vegetans syringo cystadenoma papilliferum and hidradenoma papilliferum.

Numerals in boldface type indicate the main reference to the subject

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